



NBSIR 76-1034

Chemical Thermodynamic Properties of Compounds of Sodium, Potassium and Rubidium: An Interim Tabulation of Selected Values

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Washington, D. C. 20234

April 1976

Interim Report

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no. 76-1034: of Standard Reference Data, NBS

1976

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Prepared for
Office of Standard Reference Data, NBS



U.S. DEPARTMENT OF COMMERCE, Elliot L. Richardson, Secretary
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NATIONAL BUREAU OF STANDARDS, Ernest Ambler, Acting Director

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ABSTRACT

Selected values are given for the thermochemical properties of the more common compounds of sodium and potassium. A more extensive set of selections is provided for rubidium compounds. The properties included, where data are available, are enthalpy of formation at 0 K and 298.15 K, $\Delta H_f(0)$ and $\Delta H_f(298)$, Gibbs energy of formation, entropy and heat capacity at 298.15 K; $\Delta G_f(298)$, $S(298)$ and $C_p(298)$, and the enthalpy difference between 0 K and 298.15 K, $H(298) - H(0)$. The values are consistent with the tables issued earlier in the NBS Technical Note 270 series.

Keywords: Enthalpy; entropy; Gibbs energy; heat capacity; potassium compounds; rubidium compounds; sodium compounds; standard reference data; thermochemical tables.

1. Introduction

This is an interim report on the evaluation of chemical thermodynamic properties of compounds of three of the alkali metals. It supplements the extensive set of selected values of chemical thermodynamic properties published as NBS Technical Note 270, parts 1-7. [1]. The values contained in this report are consistent with those in NBS Technical Note 270 and may be combined with them to establish the thermodynamic properties of processes.

Three tables are given here for the more common compounds of sodium and potassium and for all rubidium compounds for which data are available.

The tables contain values of the enthalpy and Gibbs energy of formation, ΔH_f° and ΔG_f° , entropy, S° , and heat capacity, C_p° , all at 298.15 K (25°C), the enthalpy of formation at 0 K, ΔH_f° , and the enthalpy difference between 0 K and 298 K, $H^\circ(298) - H^\circ(0)$. The substances included are inorganic compounds of these elements, compounds containing organic ligands that have one or two carbon atoms and their aqueous solutions.

No values are given in these tables for metal alloys or other solid solutions, fused salts, or for substances of undefined chemical composition.

The sodium and potassium tables are incomplete. They list properties of compounds evaluated in the course of preparing selections for compounds already published in the NBS Technical Note 270 series and an extensive collection of the properties of neutral electrolytes in aqueous solution at the hypothetical standard state of unit activity. The latter were calculated as the sum of the properties of the ions. Although selections will be needed for the properties of many more compounds of sodium and potassium, the selections given here should be sufficient for solution of a wide range of thermochemical problems.

The rubidium table is the complete set of selections now planned for inclusion in NBS Technical Note 270. It was prepared late in 1975. There may be a few revisions and additions based on newly reported data.

Users of these three tables are invited to comment on the selections, suggest substances for which values of properties are needed, correct errors and bring new measurements to our attention. All of these will help us improve the final tables.

2. Explanation of the contents of the tables

The following material, adapted from NBS Technical Note 270, provides definitions and conventions used in the tables.

2.1 Chemical Formulae and Physical States

The tables were reproduced from computer printout in which only capital (upper case) letters are available. Normal one-line chemical formulae are used, with the following modifications:

- Subscripts (counts of atoms) and superscripts (charge) are printed on line: $\text{NA2O}_2 = \text{Na}_2\text{O}_2$, $\text{NA}^+ = \text{Na}^+$.
- The digit "1" appears wherever necessary to separate the symbols of two chemical elements: $\text{NA1N1O}_3 = \text{NaNO}_3$.
- The centered dot, used in hydrates and minerals, is shown as a colon: $\text{NA1V1O}_4 \cdot 2\text{H}_2\text{O} = \text{NaVO}_4 \cdot 2\text{H}_2\text{O}$.
- The physical state of the substance is appended to the chemical formula in parentheses: $\text{NA1O1H(C)} = \text{NaOH}$, crystalline. The abbreviations most commonly used to denote physical states are listed in Table A. Any other notations are explained in comments within the tables.

2.2 Definition of Symbols for Thermochemical Properties

The headings used in the tables and their meanings are:

ΔH_f° , standard heat of formation at 0 K;

ΔH_f° , standard heat of formation at 298.15 K;

ΔG_f° , standard Gibbs energy (formerly free energy), G, of formation at 298.15 K;

$H_{298-H_0}^\circ = H_{298}^\circ - H_0^\circ$, enthalpy, H, of the compound in the indicated state at 298.15 K referred to its value at 0 K. If the indicated state at 298 K is gas, the corresponding state at 0 K is the hypothetical ideal gas; if the state at 298 K is solid or liquid, the corresponding state at 0 K is the thermodynamically stable crystalline solid, unless otherwise specifically indicated.

S_{298}° , standard entropy, S, at 298.15 K;

C_{298}° ; specific heat at constant pressure at 298.15 K.

The Gibbs free energy is related to the other quantities by:

$G = H - TS$, where T is the thermodynamic temperature. The enthalpy is related to the internal energy, E, by: $H = E + PV$, where P = pressure and V = volume.

All values refer to one mole of substance for the formula given.

2.3 Conventions Regarding Pure Substances

The values of the thermodynamic properties of the pure substances given in these tables are for the substances in their standard states. These standard states are defined as follows:

For a pure solid or liquid, the standard state at any temperature is the substance in the condensed phase under a pressure of one atmosphere.

For a gas the standard state at any temperature is the hypothetical ideal gas at unit fugacity, in which state the enthalpy is that of the real gas at the same temperature and at zero pressure.

The phase of a substance is indicated in parentheses at the end of the chemical formula. See section 2.1.

The values of ΔH_f° and ΔG_f° given in the tables represent the change in the appropriate thermodynamic quantity when one mole of the substance in its standard state is formed, isothermally at the indicated temperature, from the elements, each in its appropriate standard reference state. The standard reference state at 25°C for each element except phosphorus has been chosen to be the standard state that is thermodynamically stable at 25°C and at one atmosphere pressure.

For phosphorus the standard reference state is the crystalline white form; the more stable forms have not been well characterized thermo-chemically. The same reference states have been maintained for the elements at 0 K except for the liquid elements bromine and mercury for which the reference states have been chosen as the stable crystalline forms. The standard reference states for the elements are indicated in the tables by the fact that the values of ΔH_f° and ΔG_f° are exactly zero.

The values of S° represent the virtual or "thermal" entropy of the substance in the standard state at 298.15 K, omitting contributions from nuclear spins. Isotope mixing effects, etc., are also excluded except in the case of the hydrogen-deuterium (^1H - ^2H) system. Where data have been available only for a particular isotope, they have been corrected when possible to the normal isotopic composition.

The values of the enthalpies of formation of gaseous ionic species are computed on the convention that the value of ΔH_f° for the electron is zero. Conversions between 0 and 298.15 K are calculated using the value of $H_{298}^\circ - H_0^\circ = 1.481 \text{ kcal per mole of electrons}$, and assuming that the values of $H_{298}^\circ - H_0^\circ$ for the ionized and un-ionized molecules are the same.

2.4 Convention Regarding Solutions

For all dissolved substances the composition of the solvent is indicated in parentheses following the chemical formula. Except in special cases, discussed below, the number of moles of the solvent associated with one mole of solute is stated explicitly. See section 2.1 and Table A for the conventions used.

In some cases the concentration of the solute can not be specified. These are indicated as "AU" (aqueous, unspecified) for water solutions and by "U" for non-aqueous and mixed media. In all these cases the solution may be assumed to be "dilute".

The standard state for a non-dissociated solute in aqueous solution is taken as the hypothetical ideal solution of unit molality, which has been designated as "std. state, $m = 1$ ". For strong electrolytes in aqueous solution the conventional standard state is the ideal solution of unit activity (unit mean molality). For non-aqueous solutions the standard state of the solute is the hypothetical ideal solution of unit mole fraction of solute (std. state, $x_2 = 1$). The designation "A" is used for strong electrolytes in the standard state and "AO" for undissociated species in water solution. In non-aqueous media the standard state defined above is indicated by appending "X" to the formula of the solvent. The convention "std. state, $m = 1$ " is used only rarely for non-aqueous media. For it, an "M" is appended to the formula of the solvent.

The value of ΔH_f° for a solute in its standard state is equal to the apparent molal enthalpy of formation of the substance in the infinitely dilute real solution, since the enthalpy of dilution of an ideal solution is zero. At this dilution the partial molal enthalpy is equal to the apparent molal quantity. At concentrations other than the standard state, the value of ΔH_f° represents the apparent enthalpy of the reaction of formation of the solution from the elements comprising the solute, each in its standard reference state, and the appropriate total number of moles of solvent. In this representation the value of ΔH_f° for the solvent is not required. The experimental value for a heat of dilution is obtained directly as the difference between the two values of ΔH_f° at the corresponding concentrations. At finite concentrations the partial molal enthalpy of formation differs from the apparent enthalpy.

The values of the thermodynamic properties tabulated for the individual ions in aqueous solution are based on the usually convention that the values of ΔH_f° , ΔG_f° , S° and C_p° for H^+ (aq, std. state, $m = 1$) are zero. The properties of a neutral electrolyte in aqueous solution in the standard state are equal to the algebraic sum of these values for the appropriate kinds and numbers of individual ions assumed to constitute the molecule of the given electrolyte. For an ionic species e.g., HSO_4^- , the properties tabulated refer to that undissociated ion, i.e. they are not equal to the sum of those for its constituent ions. By adopting the above convention with respect to aqueous H^+ , it follows that the thermodynamic relation $\Delta H_f^\circ = \Delta H_f^\circ - T(\Delta S_f^\circ + n \cdot 0.5S^\circ (H_2))$ holds for individual ionic species, with n equal to the algebraic value of the charge. For neutral electrolytes (and gaseous ions) the normal consistency relationship applies. See section 4.

3. Unit of Energy and Fundamental Constants

All of the energy values given in these tables are expressed in terms of the thermochemical calorie. This unit, defined as equal to 4.1840 joules, is generally accepted for the presentation of chemical thermodynamic data. Values reported in other units have been converted to calories by means of the conversion factors for molecular energy given in Table B.

The following values of the fundamental physical constants have been used in these calculations:

$$R = \text{gas constant} = 8.3143 \pm 0.0012 \text{ J/deg mol} = 1.98717 \pm 0.00029 \text{ cal/deg mol}$$

$$F = \text{Faraday constant} = 96487.0 \pm 1.6 \text{ coulombs/mol} = 23060.9 \pm 0.4 \text{ cal/volt equivalent}$$

$$Z = Nhc = 11.96258 \pm 0.00107 \text{ J/cm}^{-1} \text{ mol} = 2.85912 \pm 0.00026 \text{ cal/cm}^{-1} \text{ mol}$$

$$c_2 = \text{second radiation constant} = hc/k = 1.43879 \pm 0.00015 \text{ cm deg}$$

$$0^\circ\text{C} = 273.15 \text{ K}$$

These constants are consistent with those given in the Table of General Physical Constants, recommended by the National Academy of Sciences-National Research Council [2].

These values differ slightly from those recommended by CODATA [3] in 1973. However, adoption of the CODATA recommendations would change the values in the present tables by far less than their uncertainties (see section 4.1). Thus the present values may be said to be consistent with the 1973 set of fundamental constants. The formula weights in the tables have been calculated from the molecular formula using the 1961 Table of Relative Atomic Weights based on the atomic mass of $^{12}\text{C} = 12$ exactly [4]. Values are given to four decimals places for convenience in the computer processing. Use of the more recent atomic weights [8] would produce only insignificant changes in the tabulated thermodynamic values.

4. Internal Consistency of the Tables

All of the values given in these tables have been calculated from the original articles, using consistent values for all subsidiary and auxiliary quantities. The original data were corrected where possible for differences in energy units, molecular weights, temperature scales, etc. Thus we have sought to maintain a uniform scale of energies for all the substances in the tables. In addition the tabulated values of the properties of a substance satisfy all the known physical and thermodynamic relationships among these properties. The quantities ΔH_f° , ΔG_f° , and S° at 298.15 K satisfy the relation:

$$\Delta G_f^\circ = \Delta H_f^\circ - T \Delta S_f^\circ.$$

to the precision given. The special case of solutions is discussed in section 2.4. Furthermore the calculated value of any thermodynamic quantity for a reaction is independent of the path chosen for the evaluation.

In some cases newer data may have become available on certain substances after the values were selected for these tables. Because of the need to maintain the internal consistency of the tables, it is not always possible to incorporate these newer data into the tables without a detailed analysis of the effect of such a change. Unless great care is used, relatively significant errors in calculated values of ΔH° or ΔG° for specific reactions may result from the introduction of such data. See section 4.2.

4.1 Uncertainties

The uncertainty in any value in the tables depends on the uncertainties of all the determinations in the total chain of reactions used to establish the value.

In general, when uncertainties are not stated explicitly certain rules have been followed with respect to the number of significant figures recorded. Values are tabulated such that the overall uncertainty lies between 2 and 20 units of the last figure. On the other hand, values are given so that the experimental data from which they are derived may be recovered with an accuracy equal to that of the original quantities. Thus the number of significant figures for any one value in the tables need not represent the absolute accuracy of that value. For solutions of varying composition, values are frequently tabulated to more figures to make possible the recovery of enthalpies of solution and dilution. Similarly values of ΔH_f° and $\Delta H_f^\circ_{298.15}$ may be given to different numbers of significant figures. In this instance the quantity with the lesser number of figures is used to represent the uncertainty estimate. The larger number of figures is used for the other quantity to retain the significance of the temperature correction term.

The remarks above apply directly to the tables on sodium and potassium compounds. For rubidium uncertainties are stated explicitly immediately below the value of each property. (The form is ± 0.05 , where \pm means \pm).

4.2 Relationship to Other Tables of Thermodynamic Properties

The chemical thermodynamic properties in the present table may be combined with those published in NBS Technical Note 270 [1] in order to calculate the change in a property for a process. However we recommend against these values being combined with those in any other tabulation or with a property reported in an original research paper. In particular, we warn against indiscriminate combination with the CODATA Key Values for Thermodynamics [5-7].

There are several reasons for avoiding the combination of thermochemical data from more than one table. The most important is that different large-scale tables use different thermochemical properties of formation for substances that are ubiquitous in thermochemical measurements. Outstanding examples are the common inorganic acids and their ions. Another reason is that the groups preparing different tables may have relied on different measurements as the basis for selecting property values.

It is difficult to predict a priori how a change in one selected formation property would affect values assigned to other substances because of the way these are linked by complex networks. In general, it may be expected that the advantage of internal consistency of a table will be lost if values from several sources are combined and that experimental measurements may be reproduced poorly.

No general, simple algorithm can be suggested for overcoming this problem. If it becomes necessary to extend a table of data to substances other than those tabulated, the user is advised to consult the group that prepared the table about the procedure that he plans to use.

5. Arrangement of the Tables

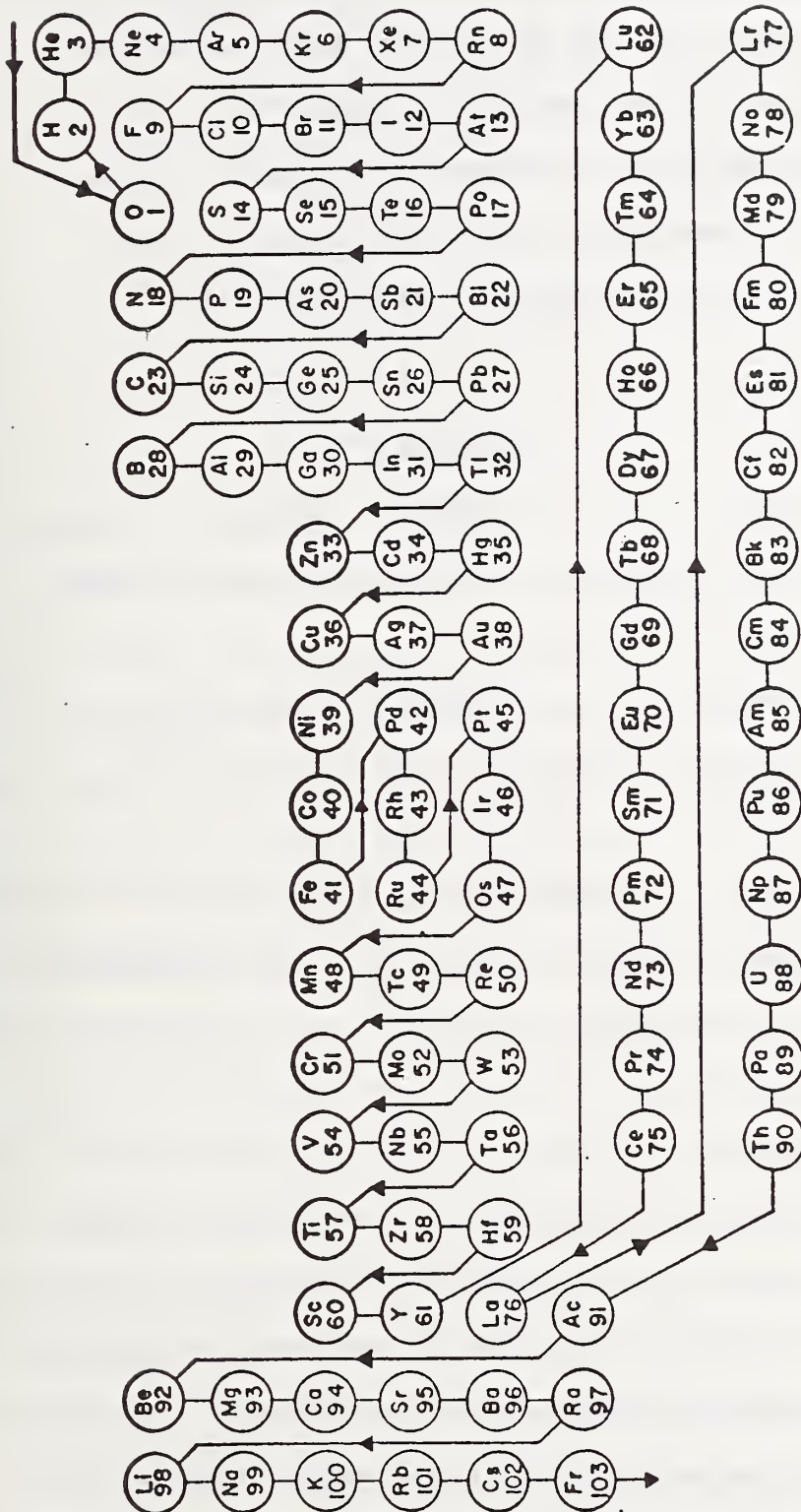
The compounds in the tables are entered according to the Standard Order of Arrangement, (see Figure 1), by the principle of latest position. In this scheme, a compound is listed under the element occurring latest in the list; water of hydration is neglected. Within a given element-table will be found all of the compounds of that element with elements occurring earlier in the order; the arrangement within a table follows the same ordering.

The selections for rubidium compounds were made late in 1975 by W. H. Evans. Those for sodium and potassium have been made by all of us and by former members of the Chemical Thermodynamics Data Center during the past decade. The extensive collection of properties of electrolytes in aqueous solution was prepared by R. H. Schumm. The tables have been printed using computer programs that are modifications of ones developed by Dr. J. B. Pedley, University of Sussex, Brighton, England, for the CATCH System of Thermochemical Tables.

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STANDARD ORDER OF ARRANGEMENT



Standard Order of Arrangement of the Elements and Compounds
based on the
Periodic Classification of the Elements

Figure 1

TABLE A: Physical State Conventions

The following conventions are used to designate the physical state of a substance. These apply to the attached table and to the NBS Thermochemical Data Bank. This information appears in a parenthetical expression appended to the molecular formula. Some of the explanations imply a thermochemical value, particularly those for solutions. These normally are used in describing enthalpy measurements.

<u>Basic Symbols</u>	<u>Explanation</u>
(G)	Gaseous, e.g., in H1CL(G) for HCl(g)
(GS)	Gaseous reference standard state for an element, e.g., O2(GS) for O ₂ (g)
(C)	Crystalline, e.g., in NH ₄ CL(C) for NH ₄ Cl(c)
(CS)	Crystalline reference standard state for an element, e.g., in RB(CS) for Rb(c)
(L)	Liquid, e.g., in H2O(L) for H ₂ O(l)
(LS)	Liquid reference standard state for an element, e.g., in BR2(LS) for Br ₂ (l)
(AM)	Amorphous
(GL)	Glassy
(A)	Hypothetical standard state of the ideal aqueous solution at unit activity. For a neutral electrolyte the value of a property is equal to the algebraic sum of the values for the ions assumed to constitute the molecule of the electrolyte, e.g. HCl(A) = H+(A) + Cl-(A). For an ionic species this notation is commonly used to refer to the undissociated ion as written. e.g. HSO ₄ ⁻ (A)

SymbolExplanation

(AO)

Hypothetical standard state of the ideal aqueous solution at unit activity of the undissociated (non-ionized) species, e.g. HF(AO), $\text{HF}_2^-(\text{AO})$.

May also be used whenever the designation (A) could be ambiguous. Note that the descriptions $\text{HSO}_4^-(\text{A})$ and $\text{HSO}_4^-(\text{AO})$ are equivalent, but that HF(A) and HF(AO) are not.

(AU)

Aqueous solution of undefined, but usually dilute, concentration, e.g. $\text{XeO}_3(\text{AU})$.

The symbols used above occasionally are modified by numbers to distinguish two substances in the same state that have the same molecular weight, as for isomers,: (AU2), (C3). They are also used in combination with descriptive material, e.g. (C:HE), (C:AL) etc. to mean "crystalline, hexagonal", "crystalline, alpha form" etc.

Special notations for substances in solutions

The notations for the "state" of a substance in solution may combine a definition of the system, e.g. HCl in 220 moles of water, and a specification of the thermochemical property associated with it. Usually the thermochemical property is the apparent integral enthalpy or free energy of formation or an absolute entropy, i.e. the formation properties of the solvent are not included. If a partial molal property is tabulated the notation D: ("D" for "differential") occurs as the first term in the state bracket. The notations given below illustrate the differences for integral and differential (partial molal) properties, and extrapolated values. Examples are given for aqueous, mixed, and non-aqueous solvents.

<u>Symbol</u>	<u>Explanation</u>
H1CL(200H2O)	An aqueous solution of specified composition, e.g. one mole of HCl in 200 moles H ₂ O. The value of ΔH _f represents the apparent integral enthalpy of formation.
H1CL(D:200H2O) and H2O(D:H1CL+200H2O)	These represent the partial molal (enthalpy) of formation of the substance in a solution of specified concentration, e.g. the partial molal enthalpy of formation of HCl and H ₂ O respectively, in a solution consisting of 1 mole HCl and 200 moles H ₂ O.
U1CL4(H1CL104+50H2O)	This describes a solute dissolved in a mixed solvent, e.g. one mole of UCl ₄ in a mixture of 1 mole of HClO ₄ and 50 moles H ₂ O. The value of ΔH _f represents the apparent integral enthalpy of formation of the substance, UCl ₄ , in the medium.
U1CL4(H1CL104+50H2O:AU)	This represents a solute at an unspecified but usually dilute concentration in a solvent mixture of fixed composition.

SymbolExplanation

U1Cl4(D:H1N1O3+50H2O) Specifies the partial molal (enthalpy) of formation of a substance in a mixed medium, e.g. the enthalpy of formation of 1 mole of UCl_4 in a large amount of a solution containing UCl_4 , HNO_3 and H_2O in the molar ratios 1:1:50.

U1Cl4(L(H1Cl):AU) Specifies a thermochemical value extrapolated from those in a particular type of solution, e.g. the integral (enthalpy) of formation of UCl_4 at an unspecified concentration in water solution obtained by extrapolation from values in solutions containing HCl at varying concentrations.

RB1I(C1H3C1N:S) Ideal solution of a substance (RbI) in a non-aqueous solvent (CH_3CN), the substance being in the standard state of unit activity on the molal scale unless otherwise indicated. (In some early entries in the Data Bank these are written ambiguously as (C1H3C1N)).

RB1I(C1H3C1N:U) Solution of a substance (RbI) in a non-aqueous solvent (CH_3CN) at an unspecified concentration. This is analogous to (AU).

RB1I(1000C1H3C1N) Solution of a substance (RbI) in a non-aqueous solvent (CH_3CN) at a specified concentration.

TABLE B
CONVERSION FACTORS FOR UNITS OF MOLECULAR ENERGY

	J/mol	cal/mol	cm ³ atm/mol	kWh/mol	Btu/lb-mol	cm ⁻¹ molecule	eV/molecule
1 J/mol =	1	2. 390057 <u>x 10⁻¹</u>	9. 86923	2. 77778 <u>x 10⁻⁷</u>	0. 429923	8. 35940 <u>x 10⁻²</u>	1. 036409 <u>x 10⁻⁵</u>
1 cal/mol =	<u>4.18400</u>	1	41.2929	1. 162222 <u>x 10⁻⁶</u>	1. 798796	3. 49757 <u>x 10⁻¹</u>	4. 33634 <u>x 10⁻⁵</u>
1 cm ³ atm/mol =	<u>0.1013250</u>	2. 42173 <u>x 10⁻²</u>	1	2. 81458 <u>x 10⁻⁸</u>	4. 35619 <u>x 10⁻²</u>	8. 47016 <u>x 10⁻³</u>	1. 050141 <u>x 10⁻⁶</u>
1 kWh/mol =	<u>3,600,000</u>	860,421	3. 55292 <u>x 10⁷</u>	1	1,547,721	300,938	37. 3107
1 Btu/lb-mol =	<u>2.32600</u>	5. 55927 <u>x 10⁻¹</u>	22. 9558	6. 46111 <u>x 10⁻⁷</u>	1	1. 944396 <u>x 10⁻¹</u>	2. 41069 <u>x 10⁻⁵</u>
1 cm ⁻¹ /molecule =	11. 96258	2. 85912	118. 0614	3. 32294 <u>x 10⁻⁶</u>	5. 14299	1	1. 239812 <u>x 10⁻⁴</u>
1 eV/molecule =	<u>96487.0</u>	23060.9	952,252	2. 68019 <u>x 10⁻²</u>	41482.0	<u>8065.73</u>	1

The underlined numbers represent the fundamental values used in deriving this table. The remaining factors were obtained by applying the relationships:

$$n_{ij} = n_{ik} \cdot n_{kj}$$

TABLES OF SELECTED VALUES OF THERMODYNAMIC PROPERTIES

Enthalpy of Formation at 0 K

Enthalpy of Formation at 298.15 K

Gibbs Energy of Formation at 298.15 K

Enthalpy at 298.15 K

Entropy at 298.15 K

Heat Capacity at 298.15 K

NBS ALKALI	METAL	CMPD.	THERMD.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL·K				
FORMULA	WT	DHO	DH298	DG298	H298 - H0	S298	CP298			
1 NA(CS)			22.9898	0	0	1.54	12.24	6.75		
2 NA(G)			22.9898	25.663	25.60	1.30	36.712	4.968		
3 NA+(A)			22.9898		-57.39	-62.593		11.1		
4 NA2O(C)		61.9790		-99.7						
5 NA2C2(C)		77.9784		-122.30						
6 NA101H1(C)			39.9972	-101.723						
7 NA1C1H(A)			39.9972	-112.36						
8 NA1C1H(2.5H2O)			39.9972	-108.100						
9 NA101H(3H2O)			39.9972		-109.053					
10 NA1C1H(4H2O)			39.9972		-110.405					
11 NA101H(4.5H2O)			39.9972	-110.847						
12 NA1C1H(5H2O)			39.9972	-111.182						
13 NA101H(6H2O)			39.9972	-111.633						
14 NA1C1H(8H2O)			39.9972	-112.071						
15 NA101H(10H2O)			39.9972	-112.248						
16 NA101H(12H2O)			39.9972	-112.326						
17 NA101H(15H2O)			39.9972	-112.370						
18 NA101H(20H2O)			39.9972	-112.380						
19 NA1C1H(25H2O)			39.9972	-112.364						
20 NA101H(30H2O)			39.9972	-112.347						
21 NA1C1H(40H2O)			39.9972	-112.316						
22 NA101H(50H2O)			39.9972	-112.293						
23 NA1C1H(75H2O)			39.9972	-112.261						
24 NA101H(100H2O)			39.9972	-112.248						
25 NA101H(150H2O)			39.9972	-112.242						
26 NA1C1H(200H2O)			39.9972	-112.239						
27 NA101H(300H2O)			39.9972	-112.244						
28 NA1C1H(500H2O)			39.9972	-112.256						
29 NA101H(800H2O)			39.9972	-112.270						
30 NA101H(1000H2O)			39.9972	-112.276						
31 NA101H1(1500H2O)			39.9972	-112.288						
32 NA101H1(2000H2O)			39.9972	-112.295						
33 NA101H1(3000H2O)			39.9972	-112.305						
34 NA101H1(5000H2O)			39.9972	-112.316						
35 NA1C1H1(7000H2O)			39.9972	-112.322						
36 NA101H1(10000H2O)			39.9972	-112.328						
37 NA101H1(20000H2O)			39.9972	-112.337						
38 NA1C1H1(50000H2O)			39.9972	-112.345						
39 NA101H1(100000H2O)			39.9972	-112.349						
40 NA101H1(00H2O)			39.9972	-112.36						

NBS ALKALI METAL CMPD.	THERMO. PROPS.	(IN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K				
		FORMULA WT	DH ₀	DG298	H298 - H0	S298
41 NA1OH:H2O(C)		58.0126	-175.557	-95.71	-78.7	19.8
42 NA1H02(A)	SCDIUM HYDROPEROXIDE: FROM Na ⁺ . HO2-	55.9966	-137.108	-136.89	-129.23	10.8
43 NA1F(C)		41.9882	-136.89	-136.89	-136.89	-14.4
44 NA1F(A)		41.9882	-136.83	-136.83	-136.83	
45 NA1F(50H2O)		41.9882	-136.798	-136.798	-136.798	
46 NA1F(75H2O)		41.9882	-136.783	-136.783	-136.783	
47 NA1F(100H2O)		41.9882	-136.775	-136.775	-136.775	
48 NA1F(150H2O)		41.9882	-136.771	-136.771	-136.771	
49 NA1F(200H2O)		41.9882	-136.775	-136.775	-136.775	
50 NA1F(300H2O)		41.9882	-136.786	-136.786	-136.786	
51 NA1F(50OH2O)		41.9882	-136.800	-136.800	-136.800	
52 NA1F(80OH2O)		41.9882	-136.806	-136.806	-136.806	
53 NA1F(100OH2O)		41.9882	-136.818	-136.818	-136.818	
54 NA1F(150OH2O)		41.9882	-136.825	-136.825	-136.825	
55 NA1F(200OH2O)		41.9882	-136.835	-136.835	-136.835	
56 NA1F(300OH2O)		41.9882	-136.846	-136.846	-136.846	
57 NA1F(500OH2O)		41.9882	-136.852	-136.852	-136.852	
58 NA1F(700OH2O)		41.9882	-136.858	-136.858	-136.858	
59 NA1F(1000OH2O)		41.9882	-136.867	-136.867	-136.867	
60 NA1F(2000OH2O)		41.9882	-136.875	-136.875	-136.875	
61 NA1F(5000OH2O)		41.9882	-136.879	-136.879	-136.879	
62 NA1F(10000OH2O)		41.9882	-136.89	-136.89	-136.89	
63 NA1F(20000OH2O)		41.9882	-212.73	-212.73	-212.73	36.2
64 NA1HF2(A)	SCDIUM BIFLUORIDE: FROM Na ⁺ , HF2-	61.9946	-200.77	-200.77	-200.77	
65 NA1CL(C)		58.4428	-98.266	-91.842	-91.842	17.23
66 NA1CL(A)		58.4428	-97.34	-93.965	-93.965	27.6
67 NA1CL(9H2O)		58.4428	-97.820	-97.820	-97.820	
68 NA1CL(10H2O)		58.4428	-97.809	-97.809	-97.809	
69 NA1CL(12H2O)		58.4428	-97.770	-97.770	-97.770	
70 NA1CL(15H2O)		58.4428	-97.707	-97.707	-97.707	
71 NA1CL(20H2O)		58.4428	-97.614	-97.614	-97.614	
72 NA1CL(25H2O)		58.4428	-97.547	-97.547	-97.547	
73 NA1CL(30H2O)		58.4428	-97.496	-97.496	-97.496	
74 NA1CL(40H2O)		58.4428	-97.425	-97.425	-97.425	
75 NA1CL(50H2O)		58.4428	-97.381	-97.381	-97.381	
76 NA1CL(75H2O)		58.4428	-97.320	-97.320	-97.320	
77 NA1CL(100H2O)		58.4428	-97.291	-97.291	-97.291	
78 NA1CL(150H2O)		58.4428	-97.266	-97.266	-97.266	
79 NA1CL(200H2O)		58.4428	-97.257	-97.257	-97.257	
80 NA1CL(300H2O)		58.4428	-97.252	-97.252	-97.252	

NBS	ALKALI	METAL	CMPD.	THERMO.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL	AND CAL/MOL•K	DH298	H298 - H0	S298
						FORMULA	WT	DH0	DH298		
81	NA1CL	(500H2O)				58•4428			-97•255		
82	NA1CL	(800H2O)				58•4428			-97•262		
83	NA1CL	(1000H2O)				58•4428			-97•267		
84	NA1CL	(1500H2O)				58•4428			-97•276		
85	NA1CL	(2000H2O)				58•4428			-97•281		
86	NA1CL	(3000H2O)				58•4428			-97•289		
87	NA1CL	(5000H2O)				58•4428			-97•298		
88	NA1CL	(7000H2O)				58•4428			-97•303		
89	NA1CL	(10000H2O)				58•4428			-97•309		
90	NA1CL	(20000H2O)				58•4428			-97•318		
91	NA1CL	(50000H2O)				58•4428			-97•326		
92	NA1CL	(100000H2O)				58•4428			-97•329		
93	NA1CL	(200000H2O)				58•4428			-97•334		
94	NA1CL	3(A)				129•3488			-91•4		
95	NA1CL	10(A)				74•4422			-83•0		
96	NA1CL	102(C)				90•4416			-73•38		
97	NA1CL	102(A)				90•4416			-73•3		
98	NA1CL	102:3H2O(C)				144•4878			-58•5		
99	NA1CL	103(C)				106•4410			-285•08		
100	NA1CL	103(A)				106•4410			-86•3		
101	NA1CL	103(6H2O)				106•4410			-86•3		
102	NA1CL	103(8H2O)				106•4410			-81•1		
103	NA1CL	103(10H2O)				106•4410			-63•4		
104	NA1CL	103(12H2O)				106•4410					
105	NA1CL	103(15H2O)				106•4410					
106	NA1CL	103(20H2O)				106•4410					
107	NA1CL	103(25H2O)				106•4410					
108	NA1CL	103(30H2O)				106•4410					
109	NA1CL	103(40H2O)				106•4410					
110	NA1CL	103(50H2O)				106•4410					
111	NA1CL	103(75H2O)				106•4410					
112	NA1CL	103(100H2O)				106•4410					
113	NA1CL	103(150H2O)				106•4410					
114	NA1CL	103(200H2O)				106•4410					
115	NA1CL	103(300H2O)				106•4410					
116	NA1CL	103(500H2O)				106•4410					
117	NA1CL	103(1000H2O)				106•4410					
118	NA1CL	103(2000H2O)				106•4410					
119	NA1CL	103(5000H2O)				106•4410					
120	NA1CL	103(10000H2O)				106•4410					

NES	ALKALI	METAL	CMPD.	THERMO.	PROPS.	(TN	270	SERIES)	IN	KCAL/MOL	AND	CAL/MOL•K
						FORMULA	WT	DHO				
1121	NA1	CL1	103	(20000H20)		106.	4410		-81.	08		
1122	NA1	CL1	103	(10000H20)		106.	4410		-81.	09		
1123	NA1	CL1	103	(GOH20)		106.	4410		-81.	1		
1124	NA1	CL1	104(C)			122.	4404		-91.	617		
1125	NA1	CL1	104(A)			122.	4404		-88.	30		
1126	NA1	CL1	104	(3.25H20)		122.	4404		-90.	48		
1127	NA1	CL1	104	(3.5H20)		122.	4404		-90.	46		
1128	NA1	CL1	104	(4H20)		122.	4404		-90.	41		
1129	NA1	CL1	104	(4.5H20)		122.	4404		-90.	36		
1130	NA1	CL1	104	(5H20)		122.	4404		-90.	31		
1131	NA1	CL1	104	(6H20)		122.	4404		-90.	19		
1132	NA1	CL1	104	(8H20)		122.	4404		-89.	98		
1133	NA1	CL1	104	(10H20)		122.	4404		-89.	81		
1134	NA1	CL1	104	(12H20)		122.	4404		-89.	66		
1135	NA1	CL1	104	(15H20)		122.	4404		-89.	48		
1136	NA1	CL1	104	(20H2C)		122.	4404		-89.	24		
1137	NA1	CL1	104	(25H20)		122.	4404		-89.	09		
1138	NA1	CL1	104	(30H20)		122.	4404		-88.	97		
1139	NA1	CL1	104	(40H20)		122.	4404		-88.	81		
1140	NA1	CL1	104	(50H20)		122.	4404		-88.	70		
1141	NA1	CL1	104	(75H20)		122.	4404		-88.	55		
1142	NA1	CL1	104	(100H20)		122.	4404		-88.	47		
1143	NA1	CL1	104	(150H20)		122.	4404		-88.	38		
1144	NA1	CL1	104	(200H20)		122.	4404		-88.	34		
1145	NA1	CL1	104	(300H20)		122.	4404		-88.	30		
1146	NA1	CL1	104	(500H20)		122.	4404		-88.	27		
1147	NA1	CL1	104	(1000H20)		122.	4404		-88.	26		
1148	NA1	CL1	104	(1500H20)		122.	4404		-88.	25		
1149	NA1	CL1	104	(2000H20)		122.	4404		-88.	26		
1150	NA1	CL1	104	(10000H20)		122.	4404		-88.	27		
1151	NA1	CL1	104	(20000H20)		122.	4404		-88.	28		
1152	NA1	CL1	104	(100000H20)		122.	4404		-88.	29		
1153	NA1	CL1	104	(100H20)		122.	4404		-88.	3		
1154	NA1	CL1	104	(10H20(C))		140.	4558		-161.	995		
1155	NA1	BR	(C)			102.	8988		-86.	296		
1156	NA1	ER	(A)			102.	8988		-86.	44		
1157	NA1	BR	(6.5H20)			102.	8988		-87.	178		
1158	NA1	BR	(8H20)			102.	8988		-87.	165		
1159	NA1	BR	(10H20)			102.	8988		-87.	099		
1160	NA1	BR	(12H20)			102.	8988		-87.	026		

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NBS	ALKALI	METAL	CMPD.	THERMO.	PROPS.	(TN	270	SERIES)	IN	KCAL/MOL	AND	CAL/MOL•K	DH298	DG298	H298	-	H0	S298	CP 298
						WT	WT		DHO		DHO	DH298							
161	NA1BR(15H20)								102.8988			-86.929							
162	NA1BR(20H20)								102.8988			-86.808							
163	NA1BR(25H20)								102.8988			-86.722							
164	NA1BR(30H20)								102.8988			-86.659							
165	NA1BR(40H20)								102.8988			-86.574							
166	NA1BR(50H20)								102.8988			-86.521							
167	NA1BR(75H20)								102.8988			-86.450							
168	NA1BR(100H20)								102.8988			-86.417							
169	NA1BR(150H20)								102.8988			-86.386							
170	NA1BR(200H20)								102.8988			-86.374							
171	NA1BR(300H20)								102.8988			-86.366							
172	NA1BR(500H20)								102.8988			-86.365							
173	NA1BR(800H20)								102.8988			-86.369							
174	NA1BR(1000H20)								102.8988			-86.372							
175	NA1BR(1500H20)								102.8988			-86.380							
176	NA1BR(2000H20)								102.8988			-86.384							
177	NA1BR(3000H20)								102.8988			-86.391							
178	NA1BR(5000H20)								102.8988			-86.400							
179	NA1BR(7000H20)								102.8988			-86.405							
180	NA1BR(10000H20)								102.8988			-86.410							
181	NA1BR(20000H20)								102.8988			-86.419							
182	NA1BR(50000H20)								102.8988			-86.426							
183	NA1BR(100000H20)								102.8988			-86.429							
184	NA1BR(00H20)								102.8988			-86.444							
185	NA1BR:2H20(C)								138.9296			-227.524							
186	NA1BR3(A)								262.7168			-88.56							
187	NA1BR5(A)								422.5348			-91.4							
188	NA1BR10(A)								118.8982			-79.9							
189	NA1BR103(C)								150.8970			-83.83							
190	NA1BR103(A)								150.8970			-77.4							
191	NA1BR103(50H20)								150.8970			-77.788							
192	NA1BR103(75H20)								150.8970			-77.647							
193	NA1BR103(100H20)								150.8970			-77.570							
194	NA1BR103(150H20)								150.8970			-77.488							
195	NA1BR103(200H20)								150.8970			-77.446							
196	NA1BR103(300H20)								150.8970			-77.406							
197	NA1BR103(500H20)								150.8970			-77.378							
198	NA1BR103(800H20)								150.8970			-77.366							
199	NA1BR103(1000H20)								150.8970			-77.364							
200	NA1BR103(1500H20)								150.8970			-77.361							

NBS ALKALI METAL CMPD.	THERMO. PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL.K	DH298	DG298	H298 - H0	S298	CP 298
		FORMULA WT	DHO	DH0	DH298			
201 NA1BR103(20000H2O)		150.8970		-77.360				
202 NA1BR103(30000H2O)		150.8970		-77.362				
203 NA1BR103(50000H2O)		150.8970		-77.367				
204 NA1BR103(70000H2O)		150.8970		-77.370				
205 NA1BR103(100000H2O)		150.8970		-77.374				
206 NA1BR103(200000H2O)		150.8970		-77.380				
207 NA1BR103(500000H2O)		150.8970		-77.386				
208 NA1BR103(1000000H2O)		150.8970		-77.390				
209 NA1BR103(000H2O)		150.8970		-77.4				
210 NA1BR2CL(A)		218.2608		-98.1	-93.3			59.02
211 NA1I(C)		149.8942		-68.780				
212 NA1I(A)		149.8942		-70.58	-74.92	4.0.7		
213 NA1I(4.5H2O)		149.8942		-71.385				
214 NA1I(5H2O)		149.8942		-71.471				
215 NA1I(6H2O)		149.8942		-71.539				
216 NA1I(6.5H2O)		149.8942		-71.544				
217 NA1I(8H2O)		149.8942		-71.504				
218 NA1I(10H2O)		149.8942		-71.408				
219 NA1I(12H2O)		149.8942		-71.314				
220 NA1I(15H2O)		149.8942		-71.193				
221 NA1I(20H2O)		149.8942		-71.042				
222 NA1I(25H2O)		149.8942		-70.938				
223 NA1I(30H2O)		149.8942		-70.863				
224 NA1I(40H2O)		149.8942		-70.764				
225 NA1I(50H2O)		149.8942		-70.702				
226 NA1I(75H2O)		149.8942		-70.619				
227 NA1I(100H2O)		149.8942		-70.579				
228 NA1I(150H2O)		149.8942		-70.541				
229 NA1I(200H2O)		149.8942		-70.525				
230 NA1I(300H2O)		149.8942		-70.512				
231 NA1I(500H2O)		149.8942		-70.508				
232 NA1I(800H2O)		149.8942		-70.511				
233 NA1I(1000H2O)		149.8942		-70.514				
234 NA1I(1500H2O)		149.8942		-70.522				
235 NA1I(2000H2O)		149.8942		-70.526				
236 NA1I(3000H2O)		149.8942		-70.532				
237 NA1I(4000H2O)		149.8942		-70.537				
238 NA1I(5000H2O)		149.8942		-70.541				
239 NA1I(7000H2O)		149.8942		-70.546				
240 NA1I(10000H2O)		149.8942		-70.550				

NBS ALKALI METAL CMPD.	THERMO. PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL.K	DH298	H298 - HO	S298
		FORMULA WT	DHO	DH298		
241 NA1I(20000H2O)		149.8942	-70.559			
242 NA1I(50000H2O)		149.8942	-70.566			
243 NA1I(100000H2O)		149.8942	-70.569			
244 NA1I(00H2O)		149.8942	-70.58			
245 NA1I:2H2O(C)		185.9250	-211.065			
246 NA1I3(A)		403.7030	-69.7	-74.9	71.3	
247 NA1I10(A)		165.8936	-83.1	-71.8	12.8	
248 NA1I103(C)		197.8924	-115.150			
249 NA1I103(A)		197.8924	-110.3	-93.2	4.2.4	
250 NA1I103(500H2O)		197.8924	-110.37			
251 NA1I103(1000H2O)		197.8924		-110.31		
252 NA1I103(2000H2O)		197.8924		-110.29		
253 NA1I103(5000H2O)		197.8924		-110.28		
254 NA1I103(100000H2O)		197.8924		-110.29		
255 NA1I103(00H2O)		197.8924		-110.3		
256 NA1I104(C)		213.8918	-101.6			
257 NA1I104(AU)	FROM NA+, 104-(AU)	213.8918	-92.6			
258 NA2I20(A)		315.7878		-144.9		
259 NA1H4I106(AU)	FRCM NA+,H4I06- (AU)	249.9226	-237.8			
260 NA1I201H(A)		29.3.8060		-117.6		
261 NA2H3I106(AU)		271.9044		-294.4		
262 NA1I1CL2(A)	FROM 2NA+,H3I06-2(AU)	220.8002		-101.1		
263 NA1I2CL(A)		312.2516	-90.3	-90.4	67.0	
264 NA1I1BR2(A)		309.7122	-92.0	-92.0		
265 NA1BR1I2(A)		356.7076	-88.0	-88.9	61.3	
266 NA1I1BR1CL(A)		265.2562		-97.6		
267 NA2S(C)		78.0436	-90.3	-86.4	19.	
268 NA2S(A)		78.0436	-106.9	-104.7	24.7	
269 NA2S(50H2O)		78.0436	-106.38			
270 NA2S(100H2O)		78.0436	-106.30			
271 NA2S(200H2O)		78.0436	-106.21			
272 NA2S(300H2O)		78.0436	-106.15			
273 NA2S(400H2O)		78.0436	-106.09			
274 NA2S(500H2O)		78.0436	-106.06			
275 NA2S(1000H2O)		78.0436	-105.94			
276 NA2S(5000H2O)		78.0436	-105.91			
277 NA2S2(A)		110.1076	-107.6	-106.2	35.0	
278 NA2S3(A)		142.1716	-108.6	-107.6	4.4.0	
279 NA2S4(A)		174.2356	-109.3	-108.7	52.9	
280 NA2S5(A)		206.2996	-109.7	-109.5	61.8	

N _b S ALKALI METAL CMPD.	THERMO. PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL·K	DH ₂₉₈	DG ₂₉₈	H ₂₉₈ - H ₀	S ₂₉₈	CP298
		FORMULA WT	DH ₀					
281	NA2S103(A)	126.0418	-266.7	-241.5	-21.			
282	NA2S104(C)	142.0412	+331.52	-303.39	35.73			30 • 42
283	NA2S104(A)	142.0412	-332.10	-303.16	33.0			-48.
284	NA2S104(28.37H ₂ O)	142.0412	-333.491					
285	NA2S104(30H ₂ O)	142.0412	-333.432					
286	NA2S104(40H ₂ O)	142.0412	-333.165					
287	NA2S104(50H ₂ O)	142.0412	-332.988					
288	NA2S104(60H ₂ O)	142.0412	-332.840					
289	NA2S104(80H ₂ O)	142.0412	+332.605					
290	NA2S104(100H ₂ O)	142.0412	-332.420					
291	NA2S104(120H ₂ O)	142.0412	-332.275					
292	NA2S104(140H ₂ O)	142.0412	-332.184					
293	NA2S104(160H ₂ O)	142.0412	-332.122					
294	NA2S104(180H ₂ O)	142.0412	-332.076					
295	NA2S104(200H ₂ O)	142.0412	-332.04					
296	NA2S104(250H ₂ O)	142.0412	-331.980					
297	NA2S104(300H ₂ O)	142.0412	-331.943					
298	NA2S104(350H ₂ O)	142.0412	-331.918					
299	NA2S104(400H ₂ O)	142.0412	-331.901					
300	NA2S104(500H ₂ O)	142.0412	-331.879					
301	NA2S104(600H ₂ O)	142.0412	-331.867					
302	NA2S104(700H ₂ O)	142.0412	-331.860					
303	NA2S104(800H ₂ O)	142.0412	-331.857					
304	NA2S104(900H ₂ O)	142.0412	-331.855					
305	NA2S104(1000H ₂ O)	142.0412	-331.855					
306	NA2S104(2000H ₂ O)	142.0412	-331.872					
307	NA2S104(3000H ₂ O)	142.0412	-331.892					
308	NA2S104(4000H ₂ O)	142.0412	-331.907					
309	NA2S104(5000H ₂ O)	142.0412	-331.92					
310	NA2S104(10000H ₂ O)	142.0412	-331.958					
311	NA2S104(20000H ₂ O)	142.0412	-331.992					
312	NA2S104(50000H ₂ O)	142.0412	-332.027					
313	NA2S104(100000H ₂ O)	142.0412	-332.047					
314	NA2S104(00H ₂ O)	142.0412	-332.10					
315	NA2S203(AU)	158.1058	-270.7					
	FROM 2N _A +	S2C3-2(AU)						
316	NA2S204(A)	174.1052	-294.9	-268.7	50.			
317	NA2S206(AU)	206.1040	-401.2					
	FROM 2N _A +	S2D6-2(AU)						
318	NA2S207(AU)	222.1034	-449.7					
	FROM 2N _A +	S2C7-2(AU)						
319	NA2S208(A)	238.1028	-434.8	-390.6	87.5			
320	NA2S306(AU)	238.1680	-401.5					
	FROM 2N _A +	S3D6-2(AU)						

NBS ALKALI METAL CMPD.	THERMO.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL	AND CAL/MOL•K	DG298	H298	- H0	S298
			FOR MUL A WT DH0	DH0	DH298				
321 NA2S4O6(AU)	FR CM 2NA+.	S4O6-2(AU)	270.2320		-407.3€				CP 298
322 NA2S5O6(AU)	FR CM 2NA+.	S5O6-2(AU)	302.2960		-410.3				
323 NA1H1S(A)				56.0618	-61.6	-59.71			29.1
324 NA1H1S1O3(A)	FR OM NA+•HS-			104.0600	-207.06	-188.74			4.7.5
325 NA1H1S1O4(A)	FR OM NA+•HSO3-			120.0594	-269.47	-243.28			45.6
	FR OM NA+•HSO4-								-9.
326 NA1H1S1O4(110H20)				120.0594	-270.16				
327 NA1H1S2O4(A)	FR OM NA+•HS2O4-			152.1234	-209.5				
328 NA1S1O3F(AU)	FR OM NA+•SO3F-(AU)			122.0504	-250.4				
329 NA2SE(A)				124.9396	-94.3				
330 NA2SE1O3(C)				172.9378	-229.43				
331 NA2SE1O3(A)				172.9378	-236.5	-213.6			31.
332 NA2SE1O4(A)				188.9372	-258.0	-230.7			41.1
333 NA1H1SE(A)	FR OM NA+•HSE-			102.9578	-53.6	-52.1			33.
334 NA1H1SE1O3(A)	FR CM NA+•HSE03-			150.9560	-180.37	-160.95			4.6.4
335 NA1H1SE1O4(A)	FR OM NA+•HSE04-			166.9554	-196.4	-170.7			49.8
336 NA2TE1O3(AU)				221.5778	-257.4				
337 NA1H5TE1O6(AU)	FR OM 2NA+•TEO3-2(AU)			251.6262	-358.9				
338 NA2F4TE1O6(AU)	FR OM NA+•H5TE06-(AU)			273.6080	-406.9				
339 NA2PO1CL6(A)	FR OM 2NA+•H4TE06-2(AU)			468.6976	-263.				
340 NA1N3(A)				65.0099	8.37	20.6			39.9
341 NA1N1O2(C)				68.9953	-85.720				
342 NA1N1O2(A)				68.9953	-82.4	-71.5			4.7.6
343 NA1N1O2(4•5H2O)				68.9953	-83.21				-12.2
344 NA1N1O2(5H2O)				68.9953	-83.19				
345 NA1N1O2(6H2O)				68.9953	-83.14				
346 NA1N1O2(8H2O)				68.9953	-83.08				
347 NA1N1O2(10H2O)				68.9953	-83.03				
348 NA1N1O2(12H2O)				68.9953	-82.99				
349 NA1N1O2(15H2O)				68.9953	-82.95				
350 NA1N1O2(20H2O)				68.9953	-82.89				
351 NA1N1O2(25H2O)				68.9953	-82.85				
352 NA1N1O2(30H2O)				68.9953	-82.82				

NBS ALKALI METAL CMPD.	THERMO. PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL•K	DH _O	DH ₂₉₈	DG ₂₉₈	H298 - H0	S298	CP 298
		FORMULA	WT						
353	NA1N1O2(4OH2O)			68.9953			-82.78		
354	NA1N1O2(5OH2O)			68.9953			-82.73		
355	NA1N1O2(75H2O)			68.9953			-82.65		
356	NA1N1O2(100H2O)			68.9953			-82.59		
357	NA1N1O2(150H2O)			68.9953			-82.50		
358	NA1N1O2(200H2O)			68.9953			-82.44		
359	NA1N1O2(300H2O)			68.9953			-82.36		
360	NA1N1O2(00H2O)			68.9953			-82.4		
361	NA1N1O3(C)			84.9947			-111.82		
362	NA1N1O3(A)			84.9947			-106.95		
363	NA1C1N1C2(AU2) SCDTUM PEROXYNITRITE; FROM NA+•, OND2-(AU)			84.9947			-87.73		
364	NA1N1O3(6H2O)			84.9947			-89.20		
365	NA1N1O3(8H2O)			84.9947			27.85		
366	NA1N1O3(10H2O)			84.9947			49.1		
367	NA1N1O3(12H2O)			84.9947			-68.6		
368	NA1N1O3(15H2O)			84.9947			-108.658		
369	NA1N1O3(20H2O)			84.9947			-108.523		
370	NA1N1O3(25H2O)			84.9947			-108.381		
371	NA1N1O3(30H2O)			84.9947			-108.253		
372	NA1N1O3(40H2O)			84.9947					
373	NA1N1O3(50H2O)			84.9947					
374	NA1N1O3(75H2O)			84.9947					
375	NA1N1O3(100H2O)			84.9947					
376	NA1N1O3(150H2O)			84.9947					
377	NA1N1O3(200H2O)			84.9947					
378	NA1N1O3(300H2O)			84.9947					
379	NA1N1O3(400H2O)			84.9947					
380	NA1N1O3(500H2O)			84.9947					
381	NA1N1O3(600H2O)			84.9947					
382	NA1N1O3(700H2O)			84.9947					
383	NA1N1O3(800H2O)			84.9947					
384	NA1N1O3(1000H2O)			84.9947					
385	NA1N1O3(3000H2O)			84.9947					
386	NA1N1O3(5000H2O)			84.9947					
387	NA1N1O3(7000H2O)			84.9947					
388	NA1N1O3(10000H2O)			84.9947					
389	NA1N1O3(20000H2O)			84.9947					
390	NA1N1O3(50000H2O)			84.9947					
391	NA1N1O3(100000H2O)			84.9947					
392	NA1N1O3(00H2C)			84.9947					

NBS ALKALI METAL CMPD.	THERMO.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL	AND CAL/MOL.K	DH298	H298 - H0	S298
			FORMULA WT	DH0	DH298			CP 298
393 NA2N202(AU)	SCDIUM HYPONITRITE		105.9918		-118.9			
394 NA1H1N202(AU)	SODIUM HYPONITRITE:	FROM NA+, HN202-(AU)	84.0100		-69.8			
395 NA1F103(AU)	SODIUM METAPHOSPHATE		101.9618		-290.9			
396 NA3P104(A)			163.9408		-477.5	-431.3		-11.
397 NA3P104(1000H2O)			163.9408		-473.6			
398 NA4F207(A)			265.9026		-772.4	-709.1		28.
399 NA4P207(2000H2O)			265.9026		-770.5			
400 NA1H2P1C2(AU)	FROM NA+, H2P02-(AU)		87.9784		-204.1			
401 NA1H2P103(AU)	FROM NA+, H2P03-(AU)		103.9778		-289.1			
402 NA1H2P103(600H2O)			103.9778		-289.1			
403 NA1H2P1C4(A)	FROM NA+, H2P04-		119.9772		-367.21	-332.76		35.7
404 NA1H2P104(400H2O)			119.9772		-366.82			
405 NA1H3P207(A)	FROM NA+, H3P207-		199.9572		-601.5	-546.2		65.
406 NA1H3P2C7(1200H2O)			199.9572		-601.5			
407 NA2H1P103(AU)	FROM 2NA+, HP03-2(AU)		125.9596		-346.3			
408 NA2H1P1C3(800H2O)			141.9590		-423.61	-385.53		20.2
409 NA2H1P104(A)	FROM 2NA+, HP04-2							
410 NA2H1P1C4(600H2O)			141.9590		-422.96			
411 NA2H2P207(A)	FROM 2NA+, H2P207-2		221.9390		-659.4	-605.7		67.
412 NA2H2P207(1200H2O)			221.9390		-659.4			
413 NA3H1P207(A)	FROM 3NA+, HP207-3		243.9208		-715.9	-659.2		53.
414 NA3H1P207(1200H2O)			243.9208		-715.9			
415 NA2P103F(A)			143.9500		-406.0			
416 NA1H1P1C3F(A)	FROM NA+, HP03F-		121.9682		-349.0			
417 NA1AS102(A)			129.9102		-159.93	-146.25		23.8
418 NA3AS104(A)			207.8886		-384.44	-342.78		3.4
419 NA1H2AS103(A)			147.9256		-228.23	-202.94		40.5
420 NA1H2AS103(400H2O)			147.9256		-228.0			
421 NA1H2AS104(A)			163.9250		-274.78	-242.63		42.
422 NA1H2AS104(3000H2O)			163.9250		-274.8			
423 NA2H1AS104(A)			185.9068		-331.40	-296.01		27.8
424 NA2H1AS104(400H2O)			185.9068		-331.2			
425 NA2AS103F(A)			187.8976		-370.78			

NES	ALKALI METAL CMPD.	THERMO. PROPS.	(IN 270 SERIES) IN KCAL/MOL AND CAL/MOL.K	
	FORMULA	WT	DHO	DG298
426	NA1H1AS1O3F(A) FR CM NA+ HASC3F-		165.9160	-316.19
427	NA1SB1O2(A)	176.7386	-143.91	
428	NA2SB2S4(A)	417.7356	-149.0	15.0
429	NA2SB2S4(400H2O)	417.7356	-167.2	
430	NA1B11CL4(A)	373.7818	-177.7	
431	NA3B11CL6(A)	490.6674	-366.29	
432	NA1B11BR4(A)	551.6058	-152.8	
433	NA2B11I4(A)	762.5772	-112.5	
434	NA2N1H4B11CL6(A)	485.7163	-322.8	
435	NA2C2(C)	70.0020	4.1	
	SCDIUM CARBIDE			
436	NA2C1O3(C)	105.9890	-270.4	32.5
437	NA2C1O3(A)	105.9890	-276.62	14.6
438	NA2C1O3(15H2O)	105.9890	-278.27	
439	NA2C1O3(20H2O)	105.9890	-278.13	
440	NA2C1O3(25H2O)	105.9890	-277.95	
441	NA2C1O3(30H2O)	105.9890	-277.78	
442	NA2C1O3(40H2O)	105.9890	-277.50	
443	NA2C1O3(50H2O)	105.9890	-277.30	
444	NA2C1O3(75H2O)	105.9890	-276.96	
445	NA2C1O3(100H2O)	105.9890	-276.76	
446	NA2C1O3(150H2O)	105.9890	-276.52	
447	NA2C1O3(200H2O)	105.9890	-276.38	
448	NA2C1O3(300H2O)	105.9890	-276.23	
449	NA2C1O3(400H2O)	105.9890	-276.15	
450	NA2C1O3(500H2O)	105.9890	-276.10	
451	NA2C1O3(600H2O)	105.9890	-276.07	
452	NA2C1O3(800H2O)	105.9890	-276.03	
453	NA2C1O3(1000H2O)	105.9890	-276.0	
454	NA2C1O3(2000H2O)	105.9890	-275.95	
455	NA2C1O3:H2O(C)	124.0044	-341.9	41.4
456	NA2C1O3:7H2O(C)	232.0968	-765.1	97.8
457	NA2C1O3:10H2O(C)	286.1430	-976.0	129.6
458	NA2C2O4(C)	133.9996	-315.0	34.
	SODIUM OXALATE			
459	NA2C2O4(A)	133.9996	-312.0	39.1
460	NA2C2O4(1000H2O)	133.9996	-311.65	
461	NA1H1C2(C)	48.0202	24.7	
	SCDIUM ACETYLIDE			
462	NA1C1H1O2(C)	68.0078	-158.02	
	SCDIUM FORMATE			
463	NA1C1H1O2(A)	68.0078	-159.10	-146.5
464	NA1C1H1C2(400H2O)	68.0078	-158.97	-9.9

CP 298
19.76
129.6
34.
36.
3.0
24.80
19.76
14.6
32.5
14.6
32.5
26.41
26.41

NBS	ALKALI	METAL	CMPD.	THERMO.	PROPS.	(TN 270	SERIES)	IN KCAL/MOL AND	CAL/MOL.K		
						FORMULA	WT	DH0	DH298	H298 - H0	S298
465	NA1C1H1O2	:2H2O(C)				104•0386		-300•7	-257•4		4•4
466	NA1C1H1O2	:3H2O(C)				122•0540		-368•4			20•94
467	NA1H1C1C3	(C)				84•0072		-226•5	-203•2		24•4
468	NA1H1C1O3	(A)				84•0072		-222•78	-202•85		35•9
469	NA1H1C1O3	(AU)				84•0072		-222•5			
470	NA1O1C1H3	(C)				54•0244				3•374	16•43
	SCDIUM	METHYLATE				54•0244		-103•63	-79•55		4•2
471	NA1O1C1H3	(A)				54•0244		-105•2			
472	NA1O1C1H3	(60C1H3O1H)				112•0178		-258•6			
473	NA1H1C2C4	(C)				112•0178		-253•0	-229•52		49•8
	SODIUM	BIOXALATE				112•0178					
474	NA1H1C2C4	(A)				112•0178		-253•3			
	FROM NA+	HC2O4-				130•0332		-330•8			
475	NA1H1C2C4	(400H2O)				82•0350		-169•41	-145•14		29•4
476	NA1H1C2C4	:H2O(C)				82•0350					19•7
477	NA1C2H3O2	(C)				82•0350		-173•55	-150•88		34•8
	SODIUM	ACETATE				82•0350					
478	NA1C2H3O2	(A)				82•0350		-170•320			
479	NA1C2H3O2	(3H2O)				82•0350					
480	NA1C2H3O2	(3•5H2O)				82•0350		-170•790			
481	NA1C2H3O2	(4H2O)				82•0350		-171•140			
482	NA1C2H3O2	(4•5H2O)				82•0350		-171•430			
483	NA1C2H3C2	(5H2O)				82•0350		-171•660			
484	NA1C2H3O2	(5•5H2O)				82•0350		-171•858			
485	NA1C2H3O2	(6H2O)				82•0350		-172•020			
486	NA1C2H3O2	(7H2O)				82•0350		-172•270			
487	NA1C2H3O2	(8H2O)				82•0350		-172•450			
488	NA1C2H3C2	(9H2O)				82•0350		-172•584			
489	NA1C2H3O2	(10H2O)				82•0350		-172•680			
490	NA1C2H3C2	(12H2O)				82•0350		-172•800			
491	NA1C2H3O2	(15H2O)				82•0350		-172•905			
492	NA1C2H3O2	(20H2O)				82•0350		-172•94			
493	NA1C2H3O2	(25H2O)				82•0350		-173•056			
494	NA1C2H3C2	(30H2O)				82•0350		-173•091			
495	NA1C2H3O2	(40H2O)				82•0350		-173•143			
496	NA1C2H3C2	(50H2O)				82•0350		-173•178			
497	NA1C2H3C2	(75H2O)				82•0350		-173•232			
498	NA1C2H3O2	(100H2O)				82•0350		-173•264			
499	NA1C2H3O2	(150H2O)				82•0350		-173•307			
500	NA1C2H3O2	(200H2O)				82•0350		-173•335			
501	NA1C2H3C2	(300H2O)				82•0350		-173•370			
502	NA1C2H3O2	(400H2O)				82•0350		-173•391			
503	NA1C2H3O2	(500H2O)				82•0350		-173•407			

			FORMULA WT	D _{HO}	DH298	DG298	H298 - HO	S298
504	NA1C2H3O2(600H20)			82.0350		-173.418		
505	NA1C2H3O2(700H20)			82.0350		-173.426		
506	NA1C2H3O2(800H20)			82.0350		-173.434		
507	NA1C2H3O2(900H20)			82.0350		-173.441		
508	NA1C2H3C2(1000H20)			82.0350		-173.446		
509	NA1C2H3O2(1500H20)			82.0350		-173.464		
510	NA1C2H3O2(2000H20)			82.0350		-173.475		
511	NA1C2H3O2(3000H20)			82.0350		-173.488		
512	NA1C2H3O2(4000H20)			82.0350		-173.496		
513	NA1C2H3C2(5000H20)			82.0350		-173.502		
514	NA1C2H3C2(7000H20)			82.0350		-173.509		
515	NA1C2H3C2(10000H20)			82.0350		-173.515		
516	NA1C2H3C2(20000H20)			82.0350		-173.525		
517	NA1C2H3O2(50000H20)			82.0350		-173.534		
518	NA1C2H3C2(100000H20)			82.0350		-173.538		
519	NA1C2H3C2(00H20)			82.0350		-173.555		
520	NA1C2H3O2:3H20(C)			136.0812		-383.19		
521	NA1C2H3O2(1100C2H501H)			82.0350		-170.8		
522	C1H201H1C101C1NA(C)			98.0344		-215.3		
	SCDIUM HYDROXYACETATE							
523	C1H201H1C1C101NA(A)			98.0344		-213.3		
524	C1H201H1C10101NA(10H20)			98.0344		-213.57		
525	C1H201H1C10101NA(15H20)			98.0344		-213.46		
526	C1H201H1C10101NA(20H20)			98.0344		-213.38		
527	C1H201H1C10101NA(25H20)			98.0344		-213.32		
528	C1H201H1C10101NA(30H20)			98.0344		-213.27		
529	C1H201H1C10101NA(40H20)			98.0344		-213.17		
530	C1H201H1C10101NA(50H20)			98.0344		-213.10		
531	C1H201H1C10101NA(100H20)			98.0344		-212.99		
532	C1H201H1C10101NA(200H20)			98.0344		-213.0		
533	C1H201H1C10101NA:1/2H20(C)			107.0421		-250.7		
534	C1H(O1H)2C1C101NA(C)			114.0338		-258.6		
	SCDIUM DIHYDROXYACETATE							
535	C1H(O1H)2C1C10101NA(200H20)			114.0338		-253.9		
536	NA101C2H5(A)			68.0516		-87.1		
537	NA101C2H5(60C2H501H)			68.0516		-111.3		
	SCDIUM ETHYLATE							
538	NA1C2H5O2(C)			84.0510		-148.4		
	SCDIUM ETHYLENE GLYCOLATE							
539	NA1C2H6O2:C1H3O1H(C)			116.0936		-211.5		
	SODIUM ETHYLENE GLYCOLATE: METHANOL							
540	C1H201H1C10101NA:C1H201H1C10101H(C)			174.0870		-377.0		
	SODIUM ACID HYDROXYACETATE							

NBS ALKALI METAL CMPD.	THERMO. PROPS.	(TN 270 SERIES) IN KCAL/MOL AND CAL/MOL•K	WT	FORMULA	DH0	DG298	H298 - H0	S298
541 NA1C2H5O2:C2H5C1H(C)								CP 298
SCDIUM ETHYLENE GLYCOLATE:ETHANOL								
542 NA1C2H5C2:C2H6O2(C)								
SCDIUM ETHYLENE GLYCOLATE:GL YCOL								
543 C1H2O1NA1C10101NA(C)								
DISODIUM HYDROXYACETATE								
544 NA1C1C1H2C1C102NA(300H2O)								
DISODIUM HYDROXYACETATE								
545 NA1C1C1H2C1C2NA:2H2O(C)								
DISODIUM HYDROXYACETATE DIHYDRATE								
546 NA1H1C1C3:NA2C103:2H2O(C)								
3NA1H1C103:NA2C103(C)								
547 NA1C2CL3O2(C)								
SCDIUM TRICHLOROACETATE								
548 NA1C2CL3O2(A)								
549 NA1C2CL3O2(C)								
550 C1CL3C1C101NA(400H2O)								
551 NA1C2CL1H2O2(A)								
SCDIUM CHLOROACETATE								
552 NA1C2CL2H1C2(A)								
SCDIUM DICHLOROACETATE								
553 NA1C2CL2H1O2(400H2O)								
554 C1H1CL2C10101NA(400H2O)								
555 NA1I:3C1H3C1H(C)								
556 NA1C2H5S1O4(600H2O)								
SODIUM ETHYLSULFATE								
557 (C1H1O)2:2NA1H1S1O3(800H2O)								
GLYOXAL SODIUM BISULFITE								
558 (C1H1O)2:2NA1H1S1O3:H2O(C)								
559 NA1C1N(C)								
((C•I•CUBIC)								
560 NA1C1N(C2)								
((C•I•ORTHCRHOMBIC)								
561 NA1C1N1O(A)								
562 NA1N1H1C1N(AU)								
563 NA1C1N:1/2H2O(C)								
564 NA1C1N:2H2O(C)								
565 NA1C1N1C(C)								
566 NA1C1N1O(A)								
567 NA1N1H1C1N(AU)								
SCDIUM CYANAMIDE								
568 NH2C10101NA(C)								
SCDIUM CARBAMATE								
569 NA1C1N1H2O2(AU)								
SODIUM SALT OF NITROMETHANE								
570 NH2C1H2C10101NA(A)								
SCDIUM SALT OF GLYCINE								

NBS ALKALI METAL CMPD. THERMO. PROPS. (TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K

FORMULA WT DH298 DG298 H298 - HO S298 CP 298

97.0497

571	NA1C2N1H4O2(Au2)	SODIUM SALT OF NITROETHANE	97.0497	-169.7	-30.06			
572	NA1C1H2C1O1O1NA(200H2O)		302.8165	23.02				
573	NA1C1N1I2(A)		201.9300	-40.75				
574	NA1(C1N1)2I(A)		81.0717					
575	NA1C1N1S(C)							
576	NA1C1N1S(A)							
577	NA1C1N1S(4H2O)							
578	NA1C1N1S(4.5H2O)							
579	NA1C1N1S(5H2O)							
580	NA1C1N1S(6H2O)							
581	NA1C1N1S(7H2O)							
582	NA1C1N1S(8H2O)							
583	NA1C1N1S(9H2O)							
584	NA1C1N1S(10H2O)							
585	NA1C1N1S(12H2O)							
586	NA1C1N1S(15H2O)							
587	NA1C1N1S(20H2O)							
588	NA1C1N1S(25H2O)							
589	NA1C1N1S(30H2O)							
590	NA1C1N1S(40H2O)							
591	NA1C1N1S(50H2O)							
592	NA1C1N1S(75H2O)							
593	NA1C1N1S(100H2O)							
594	NA1C1N1S(150H2O)							
595	NA1C1N1S(200H2O)							
596	NA1C1N1S(300H2O)							
597	NA1C1N1S(400H2O)							
598	NA1C1N1S(500H2O)							
599	NA1C1N1S(800H2O)							
600	NA1C1N1S(1000H2O)							
601	NA1C1N1S(2000H2O)							
602	NA1C1N1S(3000H2O)							
603	NA1C1N1S(4000H2O)							
604	NA1C1N1S(5000H2O)							
605	NA1C1N1S(7000H2O)							
606	NA1C1N1S(10000H2O)							
607	NA1C1N1S(20000H2O)							
608	NA1C1N1S(50000H2O)							
609	NA1C1N1S(100000H2O)							
610	NA1C1N1S(00H2O)							

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NBS ALKALI METAL CMPD.	THERMO.	PROPS.	(TN 270 SERIES) FOR MULAT	KCAL/MOL AND CAL/MOL.K	DH298	H298 - HO	S298	CP298	
611 NA1C1N1S(9C2H5O1H)			81.0717	-39.82					
612 NA1C1N1S(10C2H5O1H)			81.0717	-39.90					
613 NA1C1N1S(12C2H5O1H)			81.0717	-40.15					
614 NA1C1N1S(15C2H5O1H)			81.0717	-40.55					
615 NA1C1N1S(20C2H5O1H)			81.0717	-41.05					
616 NA1C1N1S(25C2H5O1H)			81.0717	-41.25					
617 NA1C1N1S(30C2H5O1H)			81.0717	-41.37					
618 NA1C1N1S(50C2H5O1H)			81.0717	-41.63					
619 NA1C1N1S(100C2H5O1H)			81.0717	-41.79					
620 NA1C1N1S(200C2H5O1H)			81.0717	-41.87					
621 NA1C1N1S:2S1O2(C)			209.1973	-203.4					
622 N1H2C1S1S1NA(A)			115.1517	-49.9					
CNS2H2-			137.1335	-97.4					
623 N1H1C(S2NA2)(A)									
DISODIUM DITHIOIMINOCARBONATE: FROM Na+									
2Na+ , NHCS2-2			154.1282	-477.2					
624 NA1H1S1(O1H)6(A)			188.0560	-695.4	-656.7	44.7			
625 NA2S1I1F6(C)									
626 NA2S1I1F6(A)			188.0560	-685.8	-650.9	57.4			
627 NA2S1I1F6(620H2O)			188.0560	-685.1					
628 NA1H1S1I1F6(400H2O)			166.0742	-628.0					
629 NA1S1I1CL3(A)			248.0368	-173.8	-165.4				
630 NA2S1I1CL6(AU)			377.3876	-346.7					
FRGM 2Na+ , SNCL6-2(AU)									
631 NA1SN1BR3(A)			381.4068	-147.0	-145.5				
632 NA1H1PB1O2(A)			263.1866		-143.49	74.			
FRGM Na+ , HPBO2-									
633 NA1PB1(O1H)3(A)			281.2020		-200.2				
634 NA1PB1CL3(A)			336.5388		-164.5				
635 NA1PB1BR3(A)			469.9068		-144.6				
FRGM 2Na+ , SNCL6-2(AU)									
636 NA1PB1I3(A)			610.8930		-110.1				
637 NA2PB1I4(A)			760.7872		-186.1				
638 NA2PB1P207(A)			427.1130		-605.0				
639 NA6PB1P207(AU)			519.0722		-1430.9				
640 NA6PB1(P207)2(AU)			693.0156						
FRGM 2Na+ , SNCL6-2(AU)									
641 NA1B1O2(C)			65.7996	-233.18	-220.9				
642 NA1B1O2(A)			65.7996	-241.99	-224.86	5.2			
643 NA1B1O2(220H2O)			65.7996	-242.0					
644 NA1B1O3(AU)			81.7990	-220.4					
645 NA1B1O3:4H2O(C)			153.8606	-505.3					
FRGM 2Na+ , SNCL6-2(AU)									
646 NA2B4O7(C)			201.2194	-779.98	-784.2				
647 NA2E4O7(GL)			201.2194	-779.3	-737.6	7.262	4.5-3.0	4.4-6.4	
ENTROPY IS 44.39+X							4.4-3.9	4.4-4.2	

N _B	ALKALI METAL	CMPD.	THERMC.	PROPS.	(IN 270 SERIES)	IN KCAL/MOL AND CAL/MOL·K	DG298	H298 - H0	S298	CP 298
	FORMULA	WT	DHO	DH298						
648	NA2B4O7 (A)			201.2194			-747.8			
649	NA2B4O7:4H2O (C)			273.2810			-1077.3			
650	NA2B4O7:5H2O (C)			291.2964			-1147.8			
651	NA2B4O7:10H2O (C)			381.3734			-1500.7			
652	NA2O:3B2O3 (C)			279.8396			-1097.5			
653	NA2C:3B2O3 (GL)			270.8396			-1089.1			
654	NA2C:4B2O3 (C)			340.4598			-1408.2			
655	NA2O:4B2O3 (GL)			340.4598			-1394.2			
656	NA1B1H4 (C)			37.8328	-43.100		-45.08	-29.62	3.890	24.21
657	NA1B1H4 (A)			37.8328			-45.88	-35.28		40.5
658	NA1B1H4 (35OH2O)			37.8328			-45.789			
659	NA1B1H4:2H2O (C)			73.8636			-187.6	-144.58		4.3*
660	NA1B(O1H)4 (A)			101.8304			-378.62	-338.24		38.6
661	NA1H2B1C3:H2O2 (A)			117.8298			-315.4			
662	NA1H1B4O7 (A)			179.2376			-704.4			
	FRCM NA+, HB407-									
663	NA1B1F4 (A)			109.7944			-433.8	-418.0		58.*
664	NA1B1F2(O1H)2 (A)			105.8124			-383.1			
665	NA1B1F3C1H (A)			107.8034			-422.4	-400.7		54.*
666	NA1AL1O2 (C)			81.9701			-270.96	-255.72		16.9
667	NA1AL1O2 (A)			81.9701			-277.0	-259.4		9.*
668	NA1AL1(O1H)4 (A)			1118.0009			-413.6	-372.8		42.*
669	NA3AL1F6 (C)			209.9413			-789.0	-749.7		57.0
	CRYOLITE						-775.1			
670	NA3AL1F6 (AU)			209.9413			-619.	-535.		-53.2
671	NA1AL(S1O4)2 (A)			242.0945						
	FROM NA+, 2S04-2+									
672	NA3GA1O3 (A)			186.6876			-336.			
673	NA1H2GA1O3 (A)			142.7240			-241.			
674	NA2H1GA1O3 (A)			164.7058			-289.			
	FROM 2NA+, HGA03-2									
675	NA1GA1Br4 (A)			412.3458			-215.6	-194.1		22.7
676	NA1TL1(C1N)4 (A)			331.4314				105.		
677	NA2ZN1O2 (A)			143.3484				-217.04		
678	NA1H1ZN1O2 (A)			121.3666			-171.85			
	FRCM NA+, HZNO2-									
679	NA1ZN(O1H)3 (A)			139.3820			-228.54			-169.9
680	NA2ZN(O1H)4 (A)			179.3792			-330.42			-132.3
681	NA1ZN1CL3 (A)			194.7188			-191.8			-206.5
682	NA2ZN1CL4 (A)			253.1616			-284.8			-18.4
683	NA1ZN1ER3 (A)			328.0868						-33.0
684	NA1ZN1I3 (A)			469.0730						-33.0
685	NA2ZN1I4 (A)			618.9672						
686	NA2ZN1(C1N)4 (A)			215.4212						82.*

NBS	ALKALI	METAL	CMPD.	THERMC.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL	AND CAL/MOL·K	DH298	H298 - HO	S298
						FORMULA	WT	DH0			
687	NA2ZN1	(C1N1S)4	(A)			343.6772			-73.5		
688	NA2CD1	O2	(A)			190.3784			-193.2		
689	NA1H1CD1	O2	(A)			168.3966			-149.5		
	FROM	NA+									
690	NA1CD(O1H)3	(A)				186.4120			-206.5		
691	NA2CD(O1H)4	(A)				226.4092			-306.5		
692	NA1CD1CL3	(A)				241.7488			-179.0		
693	NA1CD1BR3	(A)				375.1168			-160.0		
694	NA1CD1I3	(A)				516.1030			-124.6		
695	NA2CD1I4	(A)				665.9972			-200.7		
696	NA2CD(N3)4	(A)				326.4600			164.2		
697	NA2CD1P207	(A)				332.3230			-614.3		
698	NA1CD(C1N)3	(A)				213.4435			22.2		
699	NA2CD(C1N)4	(A)				262.4512			-3.9		
700	NA1CD(C1N1S)3	(A)				309.6355			-17.3		
701	NA1HG1CL3	(A)				329.9388			-136.5		
702	NA2HG1CL4	(A)				388.3816			-247.2		
703	NA1HG1BR3	(A)				463.3068			-127.5		
704	NA2HG1BR4	(A)				566.2056			-217.8		
705	NA1HG1I3	(A)				604.2930			-93.9		
706	NA2HG1I4	(A)				754.1872			-171.0		
707	NA1HG(C1N)3	(A)				301.6335			37.5		
708	NA2HG(C1N)4	(A)				350.6412			11.0		
709	NA1HG(C1N1S)3	(A)				397.8255			22.6		
710	NA2HG(C1N1S)4	(A)				478.8972			15.9		
711	NA2CU1O2	(A)				141.5184			-36.8		
712	NA1H1CU1O2	(A)				119.5366			-124.4		
713	NA1CU1CL2	(A)				157.4358			-120.0		
714	NA2CU1CL3	(A)				215.8786			-215.		
715	NA1CU(C1N)2	(A)				138.5656			-1.0		
716	NA2CL(C1N)3	(A)				187.5733			-28.7		
717	NA3CU(C1N)4	(A)				236.5810			-52.4		
718	NA3CU(C1N1S)4	(A)				364.8370			-93.7		
719	NA1AG1CL2	(A)				201.7658			-116.0		
720	NA1AG1BR2	(A)				290.6778			-103.8		
721	NA2AG1BR3	(A)				393.5766			-193.2		
722	NA1AG1I2	(A)				384.6686			-83.4		
723	NA2AG1I3	(A)				534.5628			-162.0		
724	NA3AG1I4	(A)				684.4570			-237.9		
725	NA1AG(C1N)2	(A)				182.8956			10.4		
726	NA1AG(S1C1N)2	(A)				247.0236			-11.2		

CP 298
DG298
DH298
CAL/MOL·K
H298 - HO
S298

62.6
105.
106.
64.
98.
76.
102.
86.
114.
67.9
101.1
137.
196.
69.4
88.7
60.

NBS ALKALI METAL CMPD. THERMO. PROPS. (IN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K

		FORMULA WT	IN KCAL/MOL AND CAL/MOL·K	DH298	H298 - HO	S298	CP298
		DHO	DH298	DH298	H298 - HO	S298	CP298
727	NA2AG(S1C1N)2(A)	328.0953	-53.3				
728	NA3AG(S1C1N)4(A)	409.1670	-94.1				
729	NA1AU1CL4(A)	361.7688	-134.4	-118.81		77.9	
730	NA1AU1BR4(A)	539.5928	-103.2	-102.6			
731	NA1AU(C1N)2(A)	271.9926	0.5	5.7		55.	
732	NA1AU(S1C1N)2(A)	336.1206	-2.4				
733	NA1AU(S1C1N)4(A)	452.2844	71.6				
734	NA2NI(C1N)4(A)	208.7612	-26.9	-12.4		80.	
735	NA3(CC(N1O2)6)(C)	403.9356	-339.6				
736	NA3(CO(N1O2)6)(14000H2O)	403.9356	-322.2				
737	NA3CO(C1N)6(A)	284.0100	97.9				
738	NA(CC(N1H3)2(N1O2)2C2O4)(C)	296.0154	-358.6				
739	NA1FE(S1O4)2(A)	270.9600	-427.0				
740	NA3FE(C1N)6(A)	280.9233	+13.5			106.9	
741	NA3FE(C1N)6(500H2O)	280.9238	-37.29				
742	NA3FE(C1N)6(800H2O)	280.9238	-37.25				
743	NA3FE(C1N)6(1000H2O)	280.9238	-37.30				
744	NA3FE(C1N)6(2000H2O)	280.9238	-37.33				
745	NA3FE(C1N)6(3000H2O)	280.9238	-37.38				
746	NA3FE(C1N)6(5000H2O)	280.9238	-37.45				
747	NA3FE(C1N)6(10000H2O)	280.9238	-37.54				
748	NA3FE(C1N)6(20000H2O)	280.9238	-37.61				
749	NA3FE(C1N)6(50000H2O)	280.9238	-37.69				
750	NA3FE(C1N)6(100000H2O)	280.9238	-37.74				
751	NA4FE(C1N)6(A)	303.9136	-120.7	-84.028	79.1		
752	NA3FE1C10(C1N)5(A)	282.9165	-119.9				
753	NA3FE1C10(C1N)5(A)	282.9165	-126.1				
754	NA3FE1C10(C1N)5(AU)	282.9165	-125.7				
755	NA3FE1C10(C1N)5(7H2O(C))	409.0243	-611.2				
756	NA1H3FE(C1N)6(A)	237.9682	51.5				
757	NA2H2FE(C1N)6(A)	259.9500	-5.9	32.18		80.	
758	NA3H1FE(C1N)6(A)	281.9318	-63.3	-27.38		84.	
759	NA1H2FE1C10(C1N)5(A)	238.9529	-11.1				
760	NA2H1FE1C10(C1N)5(A)	260.9347	-68.8				
761	NA1PD1CL3(A)	235.7488	-128.6				
762	NA2PD1CL4(A)	294.1916	-239.6	-224.8		90.	
763	NA2PD1CL6(A)	365.0976	-258.	-228.0		93.	
764	NA1PC1BR3(A)	369.1168	-111.4	-111.4			
765	NA2PD1BR4(A)	472.0156	-203.6	-201.2		98.	
766	NA2PD1BR6(A)	631.8336	-205.3	-205.3			

NBS	ALKALI	METAL	CMPD.	THERMO.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL.K		
						FORMULA	WT	DHO	H298
767	NA2PDI4(A)					659.5972		DG298	S298
768	NA2PDI6(A)					913.8060			-163.2
769	NA2FD(N1O2)4(A)					336.4016			-165.9
770	NA2FD(C1N)4(A)					256.4512			-146.3
771	NA2PDI(C1N1S)4(A)					384.7072			25.0
									-27.1
772	NA3RH1CL6(C)					384.5924			
773	NA3RH1CL6(A)					384.5924			-366.7
774	NA3RH1CL6(AU)					384.5924			-375.0
775	NA3RH1CL6:12H2O(C)					600.7772			-375.0
776	NA1RU1O4(A)					188.0574			-121.3
777	NA2RU1O4(A)					211.0472			-197.8
778	NA2RU1O4(AU)					211.0472			
	IN 0.97 NA1O1H								
779	NA1FT1CL3(A)					324.4388			-117.2
780	NA2PT1CL4(A)					382.8816			-21.3.3
781	NA2PT1CL6(C)					453.7876			
782	NA2PT1CL6(A)					453.7876			-242.
783	NA2PT1CL6(AU)					453.7876			
784	NA2PT1CL6:2H2O(C)					489.8184			-275.8
785	NA2PT1CL6:6H2O(C)					561.8800			-412.7
786	NA2PT1BR4(AU)					560.7056			-696.7
									-204.
787	NA2PT1BR6(C)					720.5236			-218.5
788	NA2PT1BR6(AU)					720.5236			
789	NA2PT1BR6:6H2O(C)					828.6160			-229.
790	NA2PT1I6(AU)					1002.4960			-647.1
791	NA2PT(C1N)4(A)					345.1412			-166.
									43.07
792	NA2R1CL6(AU)					450.8976			-262.9
793	NA3R1CL6(AU)					473.8874			-351.7
794	NA2CS1CL6(C)					448.8976			-268.
795	NA1MN1O4(A)					141.9254			-186.8
796	NA2MN1O4(A)					164.9152			-169.5
									-244.9
797	NA1MN1CL3(A)					184.2868			-271.0.8
798	NA4MN(C1N)6(A)					303.0046			-97.
799	NA1HE1O4(A)					273.1874			-245.6
800	NA2RE1CL6(A)					444.8976			-266.
801	NA2CR1O4(C)					161.9732			-294.7
									-320.8
802	NA2CR1O4(A)					161.9732			-325.38
803	NA2CR1O4(AU)					161.9732			-325.2
	IN 6000H2O + 0.1N NADH								
804	NA2CH1O4(10H2O)					161.9732			-328.1
805	NA2CR1O4(800H2O)					161.9732			-324.5
806	NA2CR1O4:4H2O(C)					234.0348			-604.2

NBS ALKALI METAL CMPD.	THERMO.	PRUPSS.	(TN 270 SERIFS)	IN KCAL/MOL AND CAL/MOL.K	DH298	DG298	H298 - H0	S298
			FORMULA	WT	DH0			
807 NA2CR104:1OH2O(C)			342•1272		-1022•7			
808 NA2CR207(C)			261•9674		-472•9			
809 NA2CR207(A)			261•9674		-471•0	-436•2		90•8
810 NA2CR207(200H2O)			261•9674		-470•5			
811 NA2CR207(300H2O)			261•9674		-470•3			
812 NA2CR207(400H2O)			261•9674		-470•1			
813 NA2CR207(500H2O)			261•9674		-470•0			
814 NA2CR207(600H2O)			261•9674		-469•9			
815 NA2CR207(800H2O)			261•9674		-469•8			
816 NA2CR207(1000H2O)			261•9674		-469•7			
817 NA2CR207(1200H2O)			261•9674		-469•6			
818 NA2CR207:2H2O(C)			297•9982		-615•4			
819 NA1H1CR104(A)			139•9914		-267•3	-245•4		58•1
FRCM Na+ • HCRO4-								
820 NA2CR104:4NA101H(C)			321•9620		-730•6			
821 NA2M0104(C)			205•9172		-350•89	-323•69		38•1
822 NA2M0104(A)			205•9172		-353•3			34•7
823 NA2M0104(AU)			205•9172		-353•1			
IN DILUTE NAOH								
824 NA2M0104:2H2O(C)			241•9480		-492•8			
825 NA2M0105(1100H2O)			221•9166		-327•5			
826 NA2M0106:H2O(C)			255•9314		-396•4			
827 NA2M0108:2H2O(C)			305•9456		-422•2			
828 NA2M0108:4H2O(C)			341•9764		-561•2			
829 NA2M0207(C)			349•8554		-536•58	-491•92		59•9
830 NA1H1M0105(1100H2O)			199•9348		-279•2			51•90
831 MO1F6:2NA1F(C)			293•9068		-660•2			
832 NA2W104(C)			293•8272		-370•2			33•40
833 NA2W104(A)			293•8272		-371•9			
834 NA2W104(AU)			293•8272		-371•6			
IN DILUTE NAOH								
835 NA2W104:2H2O(C)			329•8580		-511•5			
836 NA2W106:H2O(C)			343•8414		-418•8			
837 NA2W108:2H2O(C)			393•8556		-438•2			
838 NA2W207(C)			525•6754		-575•0	-530•0		60•8
839 NA2W4013(C)			989•3718		-993•7			
840 NASH1W6021(AU)			1555•0444		-1682•6			
841 W1F6:2NA1F(C)			381•8168		-705•3			
842 NA1V103(C)			121•9300		-273•85	-254•34		23•32
843 NA1V103(A)			121•9300		-269•7	-249•9		26•
844 NA3V104(C)			183•9090		-420•14	-391•42		45•3
845 NA3V104(A)			183•9090		-402•7			39•40
846 NA3V104:1/2H2O(C1)			192•9167		-456•4			
(C, O, H, THO)								

NBS ALKALI METAL CMPD. THERMO. PROPS. (TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K

CP298
H298 - H0 S298

		FORMULA WT	DHO	DH298	DG298	
847	NA3V104:1:2H2O(C2) (C, *PSEUDOSALT*)	192.9167	-456.6			
848	NA3V104:2H2O(C2) (C,ORTH)	219.9398	-564.1			
849	NA3V104:7/2H2O(C2) (C, *PSEUDOSALT*)	246.9629	-672.8			
850	NA3V104:7H2O(C2) (C,ORTH)	310.0168	-921.9			
851	NA3V104:8H2O(C2) (C, *PSEUDOSALT*)	328.0322	-993.0			
852	NA3V104:10H2O(C2) (C,ORTH)	364.0630	-1132.4			
853	NA3V104:10H2O(C3) (C, *PSEUDOSALT*)	364.0630	-1133.5			
854	NA3V104:12H2O(C3) (C,ORTH)	400.0938	-1272.5			
855	NA3V104:12H2O(C4) (C, *PSEUDOSALT*)	400.0938	-1273.8			
856	NA4V207(C)	305.8390	-697.62	-650.46	76.1	64.47
857	NA4V207:2H2O(C)	341.8698	-839.6			
858	NA4V207:10H2O(C)	485.9930	-1405.7			
859	NA4V207:12H2O(C)	522.0238	-1546.9			
860	NA4V207:18H2O(C)	630.1162	-1970.9			
861	NA1M2V1C4(A)	139.9454	-338.0	-306.6		4.3.
862	NA1H3V207(A)	239.8936	-508.1			
863	NA3H1V207(A)	283.8572	-616.2			
864	NA4H2V10028(A)	1051.3784	-2096.			
865	NA5H1V10028(A)	1073.3602	-2154.			
866	NA1TA1F6(A)	317.9282	-271.8			
867	NA2TA1F7(A)	359.9164	-405.4			
868	NA1MG1FE(C1N)6(A)	259.2562	-0.8			
869	NA2MG1FE(C1N)6(A)	282.2460	-73.0			
870	NA1CA1FE(C1N)6(A)	275.0242	-24.5			
871	NA2CA1FE(C1N)6(A)	298.0140	-96.6			
872	NA1SR1FE(C1N)6(A)	322.5642	-25.9			

1	K(CS)	39.1020	0	0	1.695	15.34	7.07
2	K+(A)	39.1020	-60.32	-67.70	24.5	5.2	
3	K2O(C)	94.2034	-86.4				
4	K1O1H(C)	56.1094	-101.521				
5	K1O1H(A)	56.1094	-115.29	-105.29	21.9	-30.3	
6	K1O1H(3H2O)	56.1094	-112.204				
7	K1O1H(3.5H2O)	56.1094	-112.644				
8	K1O1H(4H2O)	56.1094	-113.039				
9	K1C1H(4.5H2O)	56.1094	-113.409				
10	K1O1H(5H2O)	56.1094	-113.698				
11	K1O1H(6H2O)	56.1094	-114.028				
12	K1O1H(8H2O)	56.1094	-114.430				
13	K1O1H(10H2O)	56.1094	-114.657				
14	K1O1H(12H2O)	56.1094	-114.796				
15	K1O1H(15H2O)	56.1094	-114.920				
16	K1O1H(20H2O)	56.1094	-115.007				
17	K1O1H(25H2O)	56.1094	-115.034				
18	K1O1H(30H2O)	56.1094	-115.050				
19	K1O1H(40H2O)	56.1094	-115.072				
20	K1O1H(50H2O)	56.1094	-115.086				
21	K1O1H(75H2O)	56.1094	-115.104				
22	K1O1H(100H2O)	56.1094	-115.114				
23	K1O1H(150H2O)	56.1094	-115.129				
24	K1O1H(200H2O)	56.1094	-115.139				
25	K1O1H(300H2O)	56.1094	-115.156				
26	K1O1H(400H2O)	56.1094	-115.166				
27	K1O1H(500H2O)	56.1094	-115.175				
28	K1O1H(600H2O)	56.1094	-115.182				
29	K1O1H(700H2O)	56.1094	-115.188				
30	K1O1H(800H2O)	56.1094	-115.193				
31	K1O1H(900H2O)	56.1094	-115.198				
32	K1O1H(1000H2O)	56.1094	-115.201				
33	K1O1H(1500H2O)	56.1094	-115.215				
34	K1O1H(2000H2O)	56.1094	-115.223				
35	K1O1H(3000H2O)	56.1094	-115.234				
36	K1O1H(4000H2O)	56.1094	-115.240				
37	K1O1H(5000H2O)	56.1094	-115.246				
38	K1C1H(7000H2O)	56.1094	-115.252				
39	K1O1H(10000H2O)	56.1094	-115.258				
40	K1O1H(20000H2O)	56.1094	-115.267				

NBS ALKALI METAL CMPD. THERMO. PROPS. (TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K

CP 298
H298 - H0

	FORMULA	WT	DH	DG298	H298 - H0	S298
41 K101H(5000OH2O)	56.1094		-115.274			
42 K101H(10000OH2O)	56.1094		-115.279			
43 K101H(100H2O)	56.1094		-115.29			
44 K101H:H2O(C)	74.1248		-180.105			
45 K101H:3/2H2O(C)	83.1325		-215.262			
46 K1H102(A) POTASSIUM HYDROPEROXIDE; FROM K+, HO2-	72.1088		-98.64	-83.8	30.2	
47 K1F(C)	58.1004		-135.58	-128.53	15.91	
48 K1F(A)	58.1004		-139.82	-134.34	21.2	-20.3
49 K1F(3.5H2O)	58.1004		-138.155			
50 K1F(4H2C)	58.1004		-138.441			
51 K1F(4.5H2O)	58.1004		-138.670			
52 K1F(5H2O)	58.1004		-138.850			
53 K1F(6H2O)	58.1004		-139.101			
54 K1F(8H2C)	58.1004		-139.362			
55 K1F(10H2O)	58.1004		-139.478			
56 K1F(12H2O)	58.1004		-139.536			
57 K1F(15H2O)	58.1004		-139.579			
58 K1F(20H2O)	58.1004		-139.607			
59 K1F(25H2O)	58.1004		-139.617			
60 K1F(30H2O)	58.1004		-139.625			
61 K1F(40H2O)	58.1004		-139.633			
62 K1F(50H2O)	58.1004		-139.639			
63 K1F(75H2O)	58.1004		-139.649			
64 K1F(100H2O)	58.1004		-139.655			
65 K1F(150H2O)	58.1004		-139.666			
66 K1F(200H2O)	58.1004		-139.674			
67 K1F(300H2O)	58.1004		-139.688			
68 K1F(400H2O)	58.1004		-139.699			
69 K1F(500H2O)	58.1004		-139.707			
70 K1F(600H2O)	58.1004		-139.713			
71 K1F(700H2O)	58.1004		-139.719			
72 K1F(800H2O)	58.1004		-139.724			
73 K1F(900H2O)	58.1004		-139.728			
74 K1F(1000H2O)	58.1004		-139.731			
75 K1F(1500H2O)	58.1004		-139.745			
76 K1F(2000H2O)	58.1004		-139.753			
77 K1F(3000H2O)	58.1004		-139.764			
78 K1F(4000H2O)	58.1004		-139.771			
79 K1F(5000H2O)	58.1004		-139.776			
80 K1F(7000H2O)	58.1004		-139.782			

NBS ALKALI METAL CMPD.	THERMO. PROPS.	(TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K	DG298	H298 - H0	S298	CP298
	FOR MULAWT	DH0	DH298			
81 K1F(10000H2C)	58.1004		-139.788			
82 K1F(20000H2C)	58.1004		-139.797			
83 K1F(50000H2O)	58.1004		-139.805			
84 K1F(100000H2O)	58.1004		-139.805			
85 K1F(00H2O)	58.1004		-139.82			
86 K1F:2H2C(C)	94.1312		-278.116			
87 K1H1F2(A)	78.1068		-215.66	-205.88	46.6	
POTASSIUM EIFLUORIDE: FROM K+ + HF2-						
88 K1CL(C)	74.5550		-104.385			
89 K1CL(A)	74.5550		-100.27	-99.07	38.0	-27.4
90 K1CL(12H2O)	74.5550		-100.720			
91 K1CL(15H2O)	74.5550		-100.635			
92 K1CL(20H2O)	74.5550		-100.533			
93 K1CL(25H2O)	74.5550		-100.460			
94 K1CL(30H2O)	74.5550		-100.409			
95 K1CL(40H2O)	74.5550		-100.343			
96 K1CL(50H2O)	74.5550		-100.302			
97 K1CL(75H2O)	74.5550		-100.247			
98 K1CL(100H2O)	74.5550		-100.220			
99 K1CL(150H2O)	74.5550		-100.196			
100 K1CL(200H2O)	74.5550		-100.189			
101 K1CL(300H2O)	74.5550		-100.185			
102 K1CL(500H2O)	74.5550		-100.168			
103 K1CL(600H2O)	74.5550		-100.191			
104 K1CL(800H2O)	74.5550		-100.196			
105 K1CL(1000H2O)	74.5550		-100.200			
106 K1CL(1500H2O)	74.5550		-100.208			
107 K1CL(2000H2O)	74.5550		-100.213			
108 K1CL(3000H2O)	74.5550		-100.220			
109 K1CL(4000H2O)	74.5550		-100.225			
110 K1CL(5000H2O)	74.5550		-100.229			
111 K1CL(7000H2O)	74.5550		-100.234			
112 K1CL(10000H2O)	74.5550		-100.239			
113 K1CL(20000H2O)	74.5550		-100.248			
114 K1CL(40000H2O)	74.5550		-100.256			
115 K1CL(50000H2O)	74.5550		-100.260			
116 K1CL(00H2O)	74.5550		-100.27			
117 K1CL3(A)	145.4610			-96.5		
118 K1CL10(A)	90.5544			-76.5		
119 K1CL1C2(A)	106.5538			-63.6		
120 K1CL103(C)	122.5532			-69.645	34.17	

NBS ALKALI	METAL	CMPD.	THERMO.	PROPS.	(IN 270 SERIES)	IN KCAL/MOL AND CAL/MOL·K				
					FORMULA	WT	DH0	DH298	H298 - H0	S298
121	K1CL103(A)				122.5532		-84.0		-68.5	CP298
122	K1CL103(100H2O)				122.5532		-84.55		63.3	
123	K1CL103(150H2O)				122.5532		-84.31			
124	K1CL103(200H2O)				122.5532		-84.20			
125	K1CL103(300H2O)				122.5532		-84.09			
126	K1CL103(400H2O)				122.5532		-84.05			
127	K1CL103(500H2O)				122.5532		-84.03			
128	K1CL103(1000H2O)				122.5532		-83.99			
129	K1CL103(3000H2O)				122.5532		-83.97			
130	K1CL103(10000H2O)				122.5532		-83.98			
131	K1CL103(100000H2O)				122.5532		-83.99			
132	K1CL1C3(00H2O)				122.5532		-84.0			
133	K1CL104(C)				138.5526		-103.430		-72.46	36.1
134	K1CL104(A)				138.5526		-91.23		-69.76	68.0
135	K1CL104(600H2O)				138.5526		-91.33			
136	K1CL104(700H2O)				138.5526		-91.31			
137	K1CL104(800H2O)				138.5526		-91.30			
138	K1CL104(900H2O)				138.5526		-91.29			
139	K1CL104(1000H2O)				138.5526		-91.28			
140	K1CL104(5000H2O)				138.5526		-91.22			
141	K1CL104(10000H2O)				138.5526		-91.21			
142	K1CL104(100000H2O)				138.5526		-91.22			
143	K1CL104(00H2O)				138.5526		-91.23			
144	K1BR(C)				119.0110		-94.120			
145	K1BR(A)				119.0110		-89.37		-92.55	44.02
146	K1BR(10H2O)				119.0110		-90.195			
147	K1BR(12H2O)				119.0110		-90.084			
148	K1BR(15H2O)				119.0110		-89.954			
149	K1BR(20H2O)				119.0110		-89.860			
150	K1BR(25H2O)				119.0110		-89.713			
151	K1BR(30H2O)				119.0110		-89.643			
152	K1BR(40H2O)				119.0110		-89.549			
153	K1BR(50H2O)				119.0110		-89.490			
154	K1BR(75H2O)				119.0110		-89.407			
155	K1BR(100H2O)				119.0110		-89.368			
156	K1BR(150H2O)				119.0110		-89.332			
157	K1BR(200H2O)				119.0110		-89.317			
158	K1BR(300H2O)				119.0110		-89.306			
159	K1BR(500H2O)				119.0110		-89.303			
160	K1BR(800H2O)				119.0110		-89.305			

NBS ALKALI METAL CMPD. THERMO. PROPS. (TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K

CP298

	FORMULA	WT	DH ₀	DH ₂₉₈	DG ₂₉₈	H ₂₉₈ - H ₀	S ₂₉₈
161 K1BR(1000H20)	119.0110			-89.307			
162 K1BR(1500H20)	119.0110			-89.313			
163 K1BR(2000H20)	119.0110			-89.317			
164 K1BR(3000H20)	119.0110			-89.324			
165 K1BR(4000H20)	119.0110			-89.328			
166 K1BR(5000H20)	119.0110			-89.332			
167 K1BR(7000H20)	119.0110			-89.336			
168 K1BR(10000H20)	119.0110			-89.341			
169 K1BR(20000H20)	119.0110			-89.349			
170 K1BR(50000H20)	119.0110			-89.356			
171 K1BR(1CC000H20)	119.0110			-89.360			
172 K1BR(00H20)	119.0110			-89.37			
173 K1BR3(A)	278.8290			-91.49			
174 K1BR5(A)	438.6470			-94.3			
175 K1BR10(A)	135.0104			-82.8			
176 K1BR103(C)	167.0092			-90.130			
177 K1BR103(A)	167.0092			-80.3			
178 K1BR103(500H20)	167.0092			-80.34			
179 K1BR103(600H20)	167.0092			-80.32			
180 K1BR103(800H20)	167.0092			-80.31			
181 K1BR103(1000H20)	167.0092			-80.30			
182 K1BR103(5000H20)	167.0092			-80.27			
183 K1BR103(10000H20)	167.0092			-80.28			
184 K1BR103(100000H20)	167.0092			-80.30			
185 K1BR103(00H20)	167.0092			-80.3			
186 K1BR2CL(A)	234.3730			-101.0			
187 K1I(C)	166.0064			-78.370			
188 K1I(A)	166.0064			-73.51			
189 K1I(8H20)	166.0064			-74.848			
190 K1I(10H20)	166.0064			-74.658			
191 K1I(12H20)	166.0064			-74.509			
192 K1I(15H20)	166.0064			-74.342			
193 K1I(20H20)	166.0064			-74.149			
194 K1I(25H20)	166.0064			-74.019			
195 K1I(30H20)	166.0064			-73.927			
196 K1I(40H20)	166.0064			-73.803			
197 K1I(50H20)	166.0064			-73.724			
198 K1I(75H20)	166.0064			-73.620			
199 K1I(100H20)	166.0064			-73.565			
200 K1I(150H20)	166.0064			-73.512			

NBS ALKALI METAL CMPD.	THERMO. PROPS.	(TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K	DH298	H298 - HO	S298
		FORMULA WT	DHO	DG298	
201 K1I(200H2O)		166.0064		-73.488	
202 K1I(300H2O)		166.0064		-73.466	
203 K1I(400H2O)		166.0064		-73.457	
204 K1I(600H2O)		166.0064		-73.452	
205 K1I(800H2O)		166.0064		-73.451	
206 K1I(900H2O)		166.0064		-73.452	
207 K1I(1000H2O)		166.0064		-73.453	
208 K1I(2000H2O)		166.0064		-73.459	
209 K1I(3000H2O)		166.0064		-73.465	
210 K1I(4000H2O)		166.0064		-73.469	
211 K1I(5000H2O)		166.0064		-73.473	
212 K1I(7000H2O)		166.0064		-73.477	
213 K1I(10000H2O)		166.0064		-73.481	
214 K1I(20000H2O)		166.0064		-73.489	
215 K1I(50000H2O)		166.0064		-73.496	
216 K1I(100000H2O)		166.0064		-73.500	
217 K1I(0H2O)		166.0064		-73.51	
218 K1I ₂ (A)		419.8152		-81.7	
219 K1I10(A)		182.0058		23.2	
220 K1I103(C)		214.0046		36.20	
221 K1I103(A)		214.0046		-113.2	
222 K1I103(500H2O)		214.0046		-113.32	
223 K1I103(600H2O)		214.0046		-113.28	
224 K1I103(800H2O)		214.0046		-113.25	
225 K1I1C3(1000H2O)		214.0046		-113.23	
226 K1I103(2000H2O)		214.0046		-113.19	
227 K1I103(10000H2O)		214.0046		-113.18	
228 K1I103(100000H2O)		214.0046		-113.19	
229 K1I103(00H2O)		214.0046		-113.2	
230 K1I104(AU)		230.0040		-95.5	
FRCM K+, IC4-(AU)					
231 K2I20(A)		348.0122		-155.1	
232 K1H4I106(AU) FRCM K+, H4I06-(AU)		266.0348		-240.7	
233 K1I201H(A)		309.9182		-122.7	
234 K2H3I1C6(AU) FRCM 2K+. H3I06-2(AU)		304.1288		-300.2	
235 K1I1CL2(A)		236.9124		-106.2	
236 K1I2CL(A)		328.3638		-93.2	
237 K1I1BR2(A)		325.8244		-97.1	
238 K1PR1I2(A)		372.8196		-94.0	
239 K1I1BP1CL(A)		281.3684		-102.7	
240 K2S(A)		110.2680		-114.9	

NBS ALKALI METAL CMPD.	THERMO. PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL.K		S298
FORMULA	WT	DHO	DH298	H298 - HO	S298
241 K2S2(A)	14.2 • 3320		-113 • 4	-116 • 4	55 • 8
242 K2S3(A)	174 • 3960		-114 • 4	-117 • 8	64 • 8
243 K2S4(A)	206 • 4600		-115 • 1	-116 • 9	73 • 7
244 K2S5(A)	238 • 5240		-115 • 5	-119 • 7	82 • 6
245 K2S103(A)	158 • 2662		-272 • 5	-251 • 7	42 •
246 K2S104(C)	174 • 2656		-343 • 69	-313 • 37	53 • 8
247 K2S104(A)	174 • 2656		-337 • 96	-313 • 37	53 • 8
248 K2S2C3(AU)	190 • 3302		-276 • 5	-60 •	
FROM 2K+.	S203-2(AU)				
249 K2S204(A)	206 • 3226		-300 • 7	-278 • 9	71 •
250 K2S206(AU)	238 • 3284		-407 • 0		
FROM 2K+.	S206-2(AU)				
251 K2S207(AU)	254 • 3278		-455 • 5		
FROM 2K+.	S207-2(AU)				
252 K2S208(A)	270 • 3272		-440 • 6	-400 • 8	108 • 3
253 K2S306(AU)	270 • 3924		-407 • 3		
FROM 2K+.	S306-2(AU)				
254 K2S406(AU)	302 • 4564		-413 • 22		
FROM 2K+.	S406-2(AU)				
255 K2S506(AU)	334 • 5204		-416 • 1		
FROM 2K+.	S506-2(AU)				
256 K1H1S(C)	72 • 1740		-63 • 36		39 • 5
257 K1H1S(A)	72 • 1740		-64 • 5	-64 • 82	
FROM K+.	HS-				
258 K1H1S103(A)	120 • 1722		-209 • 99	-193 • 85	57 • 9
FROM K+.	HS03-2				
259 K1H1S104(A)	136 • 1716		-272 • 40	-248 • 39	56 • 0
FROM K+.	HS04-				
260 K1H1S204(A)	168 • 2356		-214 • 6		
FROM K+.	HS204-				
261 K1S103F(AU)	138 • 1626		-253 • 3		
FROM K+.	S03F-(AU)				
262 K2SE(A)	157 • 1640		-104 • 5		
263 K2SE103(A)	205 • 1622		-242 • 3	-223 • 8	52 • 1
FROM K+.	HS03-				
264 K2SE104(A)	221 • 1616		-263 • 8	-240 • 9	61 • 9
265 K1H1SE(A)	119 • 0700		-56 • 5	-57 • 2	43 •
FROM K+.	HSE-				
266 K1H1SE103(A)	167 • 0682		-183 • 30	-166 • 06	56 • 8
FROM K+.	HSE03-				
267 K1H1SE104(A)	183 • 0676		-199 • 3	-175 • 8	60 • 2
FROM K+.	HSE04-				
268 K2TE103(AU)	253 • 8022		-263 • 2		
FROM K+.	HSTEG6-(AU)				
269 K1H5TE1C6(AU)	267 • 7364		-361 • 8		
FROM K+.	HSTEG6-(AU)				

NBS ALKALI METAL	CMPD.	THERMO.	PROPS.	(TN 270 SERIES) FORMULA WT	IN KCAL/MOL AND DH0	DG298	H298 -	H0	S298	CP 298
270 K2H4TE106(AU)				305.8324	-412.7					
FROM 2K+ • H4TE06(AU)										
271 K2PO1CL6(A)				500.9220	5.44	-273.				50.3
272 K1N3(A)				81.1221		15.5				
FROM K+ • N3-										
273 K1NiO2(C)				85.1075	-88.490					
274 K1NiO2(A)				85.1075	-85.3	-76.6				-18.1
275 K1NiO3(C)				101.1069	-118.220					
276 K1NiO3(A)				101.1069	-109.88	-94.31				59.5
277 K10NiO2(AU2)				101.1069		-71.0				-15.5
POTASSIUM PEROXYNITRITE; FROM K+										
ON02-(AU)										
278 K1NiO3(15H2O)				101.1069		-111.679				
279 K1NiO3(20H2O)				101.1069		-111.393				
280 K1NiO3(25H2O)				101.1069		-111.187				
281 K1NiO3(30H2O)				101.1069		-111.030				
282 K1NiO3(40H2O)				101.1069		-110.811				
283 K1NiO3(50H2O)				101.1069		-110.659				
284 K1NiO3(75H2O)				101.1069		-110.427				
285 K1NiO3(100H2O)				101.1069		-110.298				
286 K1NiO3(150H2O)				101.1069		-110.154				
287 K1NiO3(200H2O)				101.1069		-110.076				
288 K1NiO3(300H2O)				101.1069		-109.994				
289 K1NiO3(400H2O)				101.1069		-109.953				
290 K1NiO3(500H2O)				101.1069		-109.928				
291 K1NiO3(600H2O)				101.1069		-109.911				
292 K1NiO3(700H2O)				101.1069		-109.900				
293 K1NiO3(800H2O)				101.1069		-109.891				
294 K1NiO3(900H2O)				101.1069		-109.884				
295 K1NiO3(1000H2O)				101.1069		-109.879				
296 K1NiO3(1500H2O)				101.1069		-109.866				
297 K1NiO3(2000H2O)				101.1069		-109.861				
298 K1NiO3(3000H2O)				101.1069		-109.856				
299 K1NiO3(5000H2O)				101.1069		-109.855				
300 K1NiO3(10000H2O)				101.1069		-109.857				
301 K1NiO3(20000H2O)				101.1069		-109.861				
302 K1NiO3(50000H2O)				101.1069		-109.866				
303 K1NiO3(100000H2O)				101.1069		-109.870				
304 K1NiO3(000H2O)				101.1069		-109.88				

NBS ALKALI METAL CMPD.	THERMO.	PROPS.	(TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K				
			FORMULA WT	DH298	DG298	H298 - HO	S298
305 K2N2O2(AU)	POTASSIUM HYONITRITE		138.2162	-124.7			CP 298
SUM 2K+ N2O2-2(AU)							
306 K1H1N2O2(AU)	POTASSIUM HYPONITRITE		100.1222	-72.7			
SUM K+ H2C2-(AU)							
307 K1P1O3(A)			118.0740	-293.8			
POTASSIUM METAFHOSPHATE							
308 K3P1O4(A)			212.2774	-486.3	-446.6	21.0	
309 K4P2O7(A)			330.3514	-784.1	-729.5	70.	
310 K1H2P1O2(A)			104.0906	-207.0			
SUM K+ H2PO2-							
311 K1H2P1O3(AU)			120.0900	-292.0			
SUM K+ H2PO3-(AU)							
312 K1H2P1O4(C)			136.0894	-375.40	-338.98	32.23	
313 K1H2P1O4(A)			136.0894	-370.14	-337.87	46.1	
SUM K+ H2PO4-							
314 K1H3P2O7(A)			216.0694	-604.4	-551.3	75.	
SUM K+ H3P2O7-							
315 K2H1F1C3(AU)			158.1840	-352.2			
SUM 2K+ HF03-2(AU)							
316 K2H1P1O4(A)			174.1834	-429.47	-395.74	41.0	
SUM 2K+ HF04-2							
317 K2H2P2O7(A)			254.1634	-665.2	-615.9	88.	
SUM 2K+ H2P2C7-2							
318 K3H1F2O7(A)			292.2574	-724.7	-674.5	84.	
SUM 3K+ HF2O7-3							
319 K2P1O3F(A)			176.1744	-416.2			
SUM K+ HF2O7-3							
320 K1H1F1O3F(A)			138.0804	-354.1			
321 K1A1S1O2(A)			146.0224	-162.86	-151.36	34.2	
322 K3A1S1O4(A)			256.2252	+393.23	-358.10	34.6	
323 K1H2A1S1O3(A)			164.0378	-231.16	-208.05	50.9	
SUM K+ H2ASO3-							
324 K1H2A1S1C4(A)			180.0372	-277.71	-247.74	52.	
SUM K+ H2ASO4-							
325 K2H1A1S1C4(A)			218.1312	-337.26	-306.22	48.6	
SUM 2K+ HASO4-2							
326 K2A1S1O3F(A)			220.1222	-380.99			
327 K1H1A1S1O3F(A)			182.0282	-321.30			
SUM K+ HASO3F-							
328 K1S8E1O2(A)			192.8508	-149.02			
329 K2S8E2S4(A)			449.9600	-173.0	-159.2	36.5	
SUM K+ H2ASO4-							
330 K1B1I1CL4(A)			389.8940	-182.8			
331 K3B1I1CL6(A)			539.0040	-381.61			
332 K1B1I1BR4(A)			567.7180	-157.9			
333 K1B1I1I4(A)			755.6996	-117.6			

NBS ALKALI METAL CMPD.	THERM.C.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL	AND CAL/MOL•K	DG298	H298 ~ H0	S298	CP 298
	FORMULA	WT	DH0	DH298	DH298	DH298	H298 ~ H0	S298	CP 298
334 K2N1H4B11CL6(A)		517.9407	138.2134	-3333.0	-3333.0	-3333.0	-3333.0	-3333.0	-3333.0
335 K2C1O3(C)	POTASSIUM CARBONATE	138.2134	138.2134	-254.7	-254.7	-254.7	-254.7	-254.7	-254.7
336 K2C1O3(A)		138.2134	138.2134	-261.57	-261.57	-261.57	-261.57	-261.57	-261.57
337 K2C1O3(25H2O)		138.2134	138.2134	-282.43	-282.43	-282.43	-282.43	-282.43	-282.43
338 K2C1O3(50H2O)		138.2134	138.2134	-282.40	-282.40	-282.40	-282.40	-282.40	-282.40
339 K2C1O3(75H2O)		138.2134	138.2134	-282.34	-282.34	-282.34	-282.34	-282.34	-282.34
340 K2C1O3(100H2O)		138.2134	138.2134	-282.29	-282.29	-282.29	-282.29	-282.29	-282.29
341 K2C1O3(150H2O)		138.2134	138.2134	-282.22	-282.22	-282.22	-282.22	-282.22	-282.22
342 K2C1O3(200H2O)		138.2134	138.2134	-282.16	-282.16	-282.16	-282.16	-282.16	-282.16
343 K2C1O3(300H2O)		138.2134	138.2134	-282.08	-282.08	-282.08	-282.08	-282.08	-282.08
344 K2C1O3(400H2O)		138.2134	138.2134	-282.02	-282.02	-282.02	-282.02	-282.02	-282.02
345 K2C1O3(500H2O)		138.2134	138.2134	-281.97	-281.97	-281.97	-281.97	-281.97	-281.97
346 K2C1O3(600H2O)		138.2134	138.2134	-281.92	-281.92	-281.92	-281.92	-281.92	-281.92
347 K2C1O3(800H2O)		138.2134	138.2134	-281.86	-281.86	-281.86	-281.86	-281.86	-281.86
348 K2C1O3(1000H2O)		138.2134	138.2134	-281.8	-281.8	-281.8	-281.8	-281.8	-281.8
349 K2C1O3(1500H2O)		138.2134	138.2134	-281.69	-281.69	-281.69	-281.69	-281.69	-281.69
350 K2C1O3(2000H2O)		138.2134	138.2134	-281.60	-281.60	-281.60	-281.60	-281.60	-281.60
251 K2C1O3:0.5H2O(C)		147.2211	147.2211	-311.4	-311.4	-311.4	-311.4	-311.4	-311.4
352 K2C1O3:1.5H2O(C)		165.23365	165.23365	-384.4	-384.4	-384.4	-384.4	-384.4	-384.4
353 K2C2O4(C)	POTASSIUM OXALATE	166.22240	166.22240	-321.3	-321.3	-321.3	-321.3	-321.3	-321.3
354 K2C2O4(A)		166.22240	166.22240	-317.8	-317.8	-317.8	-317.8	-317.8	-317.8
355 K2C2O4(25H2O)		166.22240	166.22240	-317.96	-317.96	-317.96	-317.96	-317.96	-317.96
356 K2C2O4(30H2O)		166.22240	166.22240	-317.89	-317.89	-317.89	-317.89	-317.89	-317.89
357 K2C2O4(40H2O)		166.22240	166.22240	-317.80	-317.80	-317.80	-317.80	-317.80	-317.80
358 K2C2O4(50H2O)		166.22240	166.22240	-317.73	-317.73	-317.73	-317.73	-317.73	-317.73
359 K2C2O4(75H2O)		166.22240	166.22240	-317.63	-317.63	-317.63	-317.63	-317.63	-317.63
360 K2C2O4(100H2O)		166.22240	166.22240	-317.58	-317.58	-317.58	-317.58	-317.58	-317.58
361 K2C2O4(150H2O)		166.22240	166.22240	-317.50	-317.50	-317.50	-317.50	-317.50	-317.50
362 K2C2O4(200H2O)		166.22240	166.22240	-317.52	-317.52	-317.52	-317.52	-317.52	-317.52
363 K2C2O4(400H2O)		166.22240	166.22240	-317.55	-317.55	-317.55	-317.55	-317.55	-317.55
364 K2C2O4(1000H2O)		166.22240	166.22240	-317.48	-317.48	-317.48	-317.48	-317.48	-317.48
365 K2C2O4(1500H2O)		166.22240	166.22240	-317.65	-317.65	-317.65	-317.65	-317.65	-317.65
366 K2C2O4(2000H2O)		166.22240	166.22240	-317.69	-317.69	-317.69	-317.69	-317.69	-317.69
367 K2C2O4(3000H2O)		166.22240	166.22240	-317.72	-317.72	-317.72	-317.72	-317.72	-317.72
368 K2C2O4(4000H2O)		166.22240	166.22240						
369 K2C2O4(5000H2O)		166.22240	166.22240						
370 K2C2O4(7000H2O)		166.22240	166.22240						
371 K2C2O4(10000H2O)		166.22240	166.22240						
372 K2C2O4(20000H2O)		166.22240	166.22240						
373 K2C2O4(50000H2O)		166.22240	166.22240						

NHS	ALKALI METAL CMPD.	THERMO. PROPS.	(TN 270 SERIES)	IN KCAL/MOL FORMULA WT	AND CAL/MOL/K DH0	DG298	H298 - HO	S298
374	K2C204(1000COH2O)				-317.74			CP298
375	K2C204(00H2O)				-317.8			
376	K2C204:H2O(C)				-392.8			
	POTASSIUM OXALATE (HYDRATE)							
377	K1C1H102(C)							
	POTASSIUM FORMATE							
378	K1C1H102(A)							
379	K1C1H102(400H2O)							
380	K1H1C103(C)							
	POTASSIUM BICARBONATE							
381	K1H1C103(A)							
	K+ HC03-							
382	K1H1C103(1500H2O)							
383	K101C1H3(A)							
	POTASSIUM METHYLATE							
384	K101C1H3(60C1H3O1H)							
385	K1H1C204(A)							
	POTASSIUM BIOXALATE: K+, HC204-							
386	K1H1C204(500H2O)							
387	K1C2H3O2(C)							
	POTASSIUM ACETATE							
388	K1C2H3O2(A)							
389	K1C2H3O2(4•5H2O)							
390	K1C2H3O2(5H2O)							
391	K1C2H3O2(5•5H2O)							
392	K1C2H3O2(6H2O)							
393	K1C2H3O2(7H2O)							
394	K1C2H3O2(8H2O)							
395	K1C2H3O2(9H2O)							
396	K1C2H3O2(10H2O)							
397	K1C2H3O2(12H2O)							
398	K1C2H3O2(15H2O)							
399	K1C2H3O2(20H2O)							
400	K1C2H3O2(25H2O)							
401	K1C2H3O2(30H2O)							
402	K1C2H3O2(40H2O)							
403	K1C2H3O2(50H2O)							
404	K1C2H3O2(75H2O)							
405	K1C2H3O2(100H2O)							
406	K1C2H3O2(150H2O)							
407	K1C2H3O2(200H2O)							
408	K1C2H3O2(300H2O)							
409	K1C2H3O2(400H2O)							
410	K1C2H3O2(500H2O)							

NBS ALKALI METAL CMPD. THERMO. PROPS. (TN 270 SERIES) IN KCAL/MOL AND CAL/MOL•K

F O R M U L A W T D H O D G 2 9 8 H 2 9 8 H O S 2 9 8

411	K1C2H3O2(600H2O)			98.1472		-176.344		
412	K1C2H3O2(800H2O)			98.1472		-176.361		
413	K1C2H3O2(900H2O)			98.1472		-176.368		
414	K1C2H3O2(1000H2O)			98.1472		-176.373		
415	K1C2H3O2(1500H2O)			98.1472		-176.393		
416	K1C2H3O2(2000H2O)			98.1472		-176.404		
417	K1C2H3O2(3000H2O)			98.1472		-176.417		
418	K1C2H3O2(4000H2O)			98.1472		-176.425		
419	K1C2H3O2(5000H2O)			98.1472		-176.431		
420	K1C2H3O2(7000H2O)			98.1472		-176.438		
421	K1C2H3O2(10000H2O)			98.1472		-176.445		
422	K1C2H3O2(20000H2O)			98.1472		-176.455		
423	K1C2H3O2(50000H2O)			98.1472		-176.464		
424	K1C2H3O2(100000H2O)			98.1472		-176.468		
425	K1C2H3O2(00H2O)			98.1472		-176.48		
426	C1H2O1HIC1O101K(A)			114.1466		-217.5		
	POTASSIUM HYDROXYACETATE							
427	K1C1H2O1H1C1O10(A)			114.1466		-216.2		
428	C1H2O1H1C1O101K(200H2O)			114.1466		-216.0		
429	C1H2O1H1C1O101K:1/2H2O(C)			123.1543		-254.7		
	POTASSIUM HYDROXYACETATE HEMIHYDRATE							
430	K1O1C1H2C1H3(A)			84.1638		-92.2		
	POTASSIUM ETHYLATE							
431	K1O1C1H2C1H3(60C2H5O1H)			84.1638		-115.9		
432	S(K2C1O3:3/2H2O):4K1H1C1O3(C)			1226.6601		-2840.		
423	K1C2CL3O2(A)			201.4822		-183.7		
	POTASSIUM TRICHLOROACETATE							
434	K1C2CL3C2(50H2O)			201.4822		-183.8		
435	K1C2CL1H2O2(A)			132.5922		-180.13		
	POTASSIUM CHLOROACETATE							
436	K1C2CL2H1O2(A)			167.0372		-182.7		
	POTASSIUM DICHLOROACETATE							
437	(C1H1O)2:2K1H1S1O3(C)			298.3816		-531.1		
	GLYXAL POTASSIUM BISULFITE							
438	(C1H1O)2:2K1H1S1O3(800H2O)			298.3816		-518.2		
439	K1C1N(C)			65.1199		-27.10		
	{C, I, CUBIC}							
440	K1C1N(A)			65.1199		-24.3		
						-26.5		4.7e0
441	K1C1N(200H2O)			65.1199		-24.0		
442	K1C1N1O(C)			81.1193		-100.04		
	POTASSIUM CYANATE							
443	K1C1N1O(A)			81.1193		-95.2		
444	K1C1H2N1O2(AU)			99.1347		-80.6		50.0
	POTASSIUM SALT OF NITROMETHANE: K+							
	CH2NO2-							

NBS ALKALI METAL CMFD.	THERMO.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL·K				
			FORMULA	WT	DHO	DH298	H298 - H0	S 298
445 K1O1C1C1C1O1N1H2(C) POTASSIUM CXAMATE			127.1453		-220.05			
446 K1C2N1H4O2(A) POTASSIUM SALT OF GLYCINE: K+.			113.1619		-172.60	-142.98		53.0
447 K1C2N1H4O2(A2) POTASSIUM SALT OF NITROETHANE: K+.			113.1619		-90.6			
C2NH4O2-								
448 K1C1N1I2(A) K+.			318.9287		-35.17			
CN12-								
449 K1I(C1N)2(A) K+.			218.0422		17.91			
{CN)2I-								
K1C1N1S(C) POTASSIUM THIOCYANATE			97.1839		-47.84	-40.70		20.6
451 K1C1N1S(A)			97.1839		-42.05	-45.55		59.0
452 K1C1N1S(2H2O)			97.1839		-44.690			-4.4
453 K1C1N1S(2•5H2O)			97.1839		-44.560			
454 K1C1N1S(3H2O)			97.1839		-44.440			
K1C1N1S(4H2O)			97.1839		-44.220			
456 K1C1N1S(4•5H2O)			97.1839		-44.112			
457 K1C1N1S(5H2O)			97.1839		-44.010			
458 K1C1N1S(6H2O)			97.1839		-43.840			
459 K1C1N1S(7H2O)			97.1839		-43.688			
K1C1N1S(8H2O)			97.1839		-43.570			
460 K1C1N1S(9H2O)			97.1839		-43.458			
461 K1C1N1S(10H2O)			97.1839		-43.360			
462 K1C1N1S(12H2O)			97.1839		-43.195			
463 K1C1N1S(15H2O)			97.1839		-43.015			
K1C1N1S(18H2O)			97.1839		-42.805			
465 K1C1N1S(20H2O)			97.1839		-42.661			
466 K1C1N1S(25H2O)			97.1839		-42.557			
467 K1C1N1S(30H2O)			97.1839		-42.418			
468 K1C1N1S(40H2O)			97.1839		-42.330			
469 K1C1N1S(50H2O)			97.1839		-42.015			
K1C1N1S(75H2O)			97.1839		-42.204			
470 K1C1N1S(100H2O)			97.1839		-42.139			
471 K1C1N1S(150H2O)			97.1839		-42.074			
472 K1C1N1S(200H2O)			97.1839		-42.043			
473 K1C1N1S(300H2O)			97.1839		-42.001			
474 K1C1N1S(400H2O)			97.1839		-42.005			
K1C1N1S(500H2O)			97.1839		-42.010			
475 K1C1N1S(600H2O)			97.1839		-42.003			
476 K1C1N1S(700H2O)			97.1839		-41.998			
477 K1C1N1S(1000H2O)			97.1839		-41.994			
478 K1C1N1S(1500H2O)			97.1839		-41.998			
479 K1C1N1S(2000H2O)			97.1839		-42.001			
K1C1N1S(3000H2O)			97.1839		-42.005			
480 K1C1N1S(4000H2O)			97.1839		-42.010			
481 K1C1N1S(4000H2O)			97.1839		-42.010			

NBS ALKALI METAL CMPD. THERMO. PROPS. (TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K	FORMULA	WT	DHO	DH298	H298 - H0	S298	CP298
482 K1C1N1S(5000H2O)				97.1839	-42.012		
483 K1C1N1S(7000H2O)				97.1839	-42.017		
484 K1C1N1S(10000H2O)				97.1839	-42.021		
485 K1C1N1S(20000H2O)				97.1839	-42.028		
486 K1C1N1S(5000H2O)				97.1839	-42.036		
487 K1C1N1S(100000H2O)				97.1839	-42.039		
488 K1C1N1S(00H2O)				97.1839	-42.05		
489 K1C1N1S:0.5S102(C)				129.2153	-88.9	-77.0	35.7
490 K1C1N1S:2S102(C)				225.3095	-209.8	-183.2	71.2
491 K1C1N1S2H2(A)				131.2639	-52.8		
POTASSIUM DITHIOCARBAMATE: K+, CNS2H2-							
492 K2C1N1S2H(A)				169.3579	-103.2		
DIPOTASSIUM DITHIOCIMINOCARBONATE: 2K+							
CNS2H2							
493 K1H1S1(01H)6(A)				170.2404	-482.3		
494 K2S11F6(C)				220.2804	-709.		
495 K2S11F6(A)				220.2804	-691.6	-661.1	78.2
496 K1S1N1CL3(A)				264.1510	-176.7	-170.5	87.
497 K2SN1CL6(C)				409.6120	-355.6		
498 K2SN1CL6(AU)				409.6120	-352.5		
FROM 2K+ SNCL6-2(AU)							
499 K1SN1BR3(A)				397.5190	-149.9	-150.6	85.
500 K1H1PB102(A)				279.2988	-148.60	-148.60	
501 K1PB1(01H)3(A)				297.3142	-205.3	-205.3	
502 K1PB1CL3(A)				352.6510	-169.6		
503 K1PB1BR3(A)				486.0190	-149.7		
504 K1PB1I3(A)				627.0052	-115.2		
505 K2PB1I4(A)				793.0116	-196.3		
506 K2PB1P2C7(A)				459.3374	-615.2		
507 K6PB1P207(AU)				615.7454	-1448.5		
508 K1B1O2(A)				81.9118	+244.92	-229.97	15.6
509 K2B4O7(A)				233.4438	-1422.3	-758.0	
510 K2O:4B2C3(C)				372.6842	-1405.0		
511 K2O:4B2C3(GL)				372.6842			
K+, HB4O7-							
512 K1B1H4(C)				53.9450	-54.21	-38.21	25.48
513 K1B1H4(A)				53.9450	-48.81	-40.39	50.9
514 K1B1(01H)4(A)				117.9426	-381.55	-343.35	49.0
515 K1H2B1O3:H2O2(A)				133.9420	-320.5	-320.5	
FROM K+, H2B03:H2O2-							
516 K1H1B4O7(A)				195.3498	-709.5		
K+, HB4O7-							
517 K1B1F4(C)				125.9066	-448.69	-449.8	426.8
518 K1B1F4(A)				125.9066	-436.7	-423.1	26.4
519 K1B1F2(C1H)2(A)				121.9246	-388.2	-388.2	68.

NBS ALKALI	METAL	CMPD.	THERMO.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL·K	FORMULA WT	DHO	DG298	H298 - H0	S298
520	K1B1F3O1H(C)				123.9156	-432.9					CP 298
521	K1B1F3O1H(A)				123.9156	-425.3					
522	K1B1F3O1H(400H2O)				123.9156	-425.6					64.
523	K2B3F4O3O1H(C)				251.6362	-893.1					
524	K2B3F4O3O1H(800H2O)				251.6362	-874.1					
525	K1AL1O2(A)				98.0823	-279.9					20.
526	K1AL(C1H)4(A)				134.1131	-416.5					53.
527	K3AL1F6(AU)				258.2779	-783.9					
528	K1AL(S1O4)2(C)				258.2067	-590.38					46.12
529	K1AL(S1C4)2(A)				258.2067	-622.					48.9
	FROM AL+3+, K+, 2S042-										-42.8
530	K1AL(S1O4)2(AU)				258.2067	-619.9					
531	K1AL(S1C4)2:H2O(C)				276.2222	-668.7					
532	K1AL(S1C4)2:2H2O(C)				294.2375	-743.					
533	K1AL(S1C4)2:3H2O(C)				312.2529	-815.					
534	K1AL(S1C4)2:12H2O(C)				474.3915	-1448.88					155.6
535	K3GA1O3(A)				235.0242	-351.					
536	K1H2GA1O3(A)				158.8362	-246.					
	FRCM K+· H2GA03										
537	K2H1GA1O3(A)				196.9302	-299.					
	FROM 2K+. HGAC3-2										
538	K1GA1BR4(A)				428.4580	-218.5					
539	K1TL(C1N)4(A)				347.5436	100.					
540	K2ZN1O2(A)				175.5728	-227.25					
541	K1H1ZN1C2(A)				137.4788	-176.96					
542	FROM K+· HZNO2-										
	K1ZN1O3H3(A)										
543	K2ZN(O1H)4(A)				155.4942	-233.65					
544	K1ZN1CL3(A)				211.6036	-340.63					
					210.8310	-196.9					
545	K2ZN1CL4(A)				285.3860	-294.6					
546	K1ZN1BR3(A)				344.1990	-175.0					
547	K1ZN1I3(A)				485.1852	-137.4					
548	K2ZN1I4(A)				651.1916	-216.7					
549	K2ZN(C1N)4(A)				247.6456	-38.8					103.
550	K2ZN(C1N)4(600H2O)				247.6456	-38.8					
551	K2ZN(C1NIS)4(A)				375.9016	-83.7					
552	K2CD1O2(A)				222.6028	-203.4					
553	K1H1CD1C2(A)				184.5088	-154.6					
	FRCM K+· HCD02-										
554	K1CD(O1H)3(A)				202.5242	-211.6					
555	K2CD(O1H)4(A)				258.6336	-316.7					
556	K1CD1CL3(A)				257.8610	-184.1					
557	K1CD1BR3(A)				391.2290	-165.1					
558	K1CD1I3(A)				532.2152	-129.7					

NBS ALKALI METAL CMPD. THERMC. PROPS. (TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K

CP 298

	FORMULA	WT	DH	DH298	DG298	H298 - H0	S298
559 K2CD1I4(A)				698.22216	-202.3	-210.9	127.
560 K1CD(N3)3(A)				277.5623		159.1	
561 K2CD(N3)4(A)				358.6844		174.1	
562 K2CD1P2O7(A)				364.5474		-624.5	
563 K1CD(C1N)3(A)				229.5557		17.1	
564 K2CD(C1N)4(A)				294.6756	-18.3	-14.1	
565 K2CD(C1N)4(8000H2O)				294.6756	-18.3		
566 K1CDC1N1S)3(A)				325.7477		-22.4	
567 K1HG1CL3(A)				346.0510	-153.2	-141.6	75.
568 K2HG1CL4(A)				420.6060	-253.0	-242.2	119.
569 K1HG1BR3(A)				479.4190	-130.4	-129.7	86.
570 K2HG1BR4(A)				598.4300	-223.6	-224.1	123.
571 K1HG1I3(A)				620.4052	-96.8	-103.2	97.
572 K2HG1I4(A)				786.4116	-176.8	-186.0	135.
573 K1HG(C1N)3(A)				317.7457	34.6	43.0	78.3
574 K2HG(C1N)4(A)				382.8656	5.2	12.4	122.
575 K1HG(C1N1S)3(A)				413.9377		10.8	
576 K2HG(C1N1S)4(A)				511.1216	-42.6	-37.1	158.
577 K2CU1O2(A)				173.7428		-179.3	
578 K1H1CU1C2(A)				135.6488		-129.5	
579 K1CL1CL2(A)				173.5480		-125.1	
580 K1CU1CL3(A)				209.0010		-225.	
581 K1CU(C1N)2(A)				154.6778		-6.1	
582 K2CU(C1N)3(A)				219.7977		-38.9	
583 K3CU(C1N)4(A)				284.9176		-67.7	
584 K3CU(C1N1S)4(A)				413.1736	-102.5	-116.1	227.
585 K1AG1CL2(A)				217.8780	-118.9	-119.2	75.8
586 K1AG1BR2(A)				306.7900		-108.9	
587 K2AG1BR3(A)				425.8010		-203.4	
588 K1AG1I2(A)				400.7808		-88.5	
589 K2AG1I3(A)				566.7872	-164.1	-172.2	109.5
590 K3AG1I4(A)				732.7936		-253.2	
591 K1AG(C1N)2(A)				199.0078	4.3	5.3	71.
592 K1AG(S1C1N)2(A)				263.1358		-16.3	
593 K2AG(S1C1N)3(A)				360.3197		-63.5	
594 K3AG(S1C1N)4(A)				457.5036		-109.4	
595 K1AU1CL4(A)				377.8810	-137.3	-123.92	88.3
596 K1AU1BR4(A)				555.7050	-106.1	-107.7	104.8
597 K1AU(C1N)2(A)				288.1048	-2.4	0.6	65.
598 K1AL(S1C1N)2(A)				352.2328		-7.5	

NBS ALKALI METAL CMPD.	THEPMG.	PROPS.	(TN 270 SERIES)	IN K CAL/MOL AND CAL/MOL·K	DG298	H298 - H0	S298	CP298
			FORMULA WT	DHO				
599 K1AUS1C1N)4(A)			4 68. 3966		-32.0	66.5		
600 K2N1(C1N)4(A)			240. 9856		-231.1	-22.6		101.
601 K(CC(N1)3)2(N1O2)4(C)			316. 1186		-217.9			
602 K(CC(N1)3)2(N1O2)4(300COH2O)			316. 1186		-752.2			
603 K3(CO(C2O4)3)(20000H2O)			440. 2992					
604 K3(CO(C2O4)3):3H2O(C)			494. 3454		-975.0	96.0		
605 K3CC(C1N)6(C)			332. 3466		332. 3466	129.1		
606 K3CC(C1N)6(A)			332. 3466		332. 3466			
607 K(CC(N1)3)2(N1O2)2C2O4(C)			312. 1276		-366.2			
608 K(CC(N1)3)2(N1O2)2C2O4(17000H2O)			312. 1276		-356.4			
609 K1FE(S1O4)2(A)			287. 0722		-432.1			
610 K1FE(C1N)6-2(AO)			251. 0564		251. 0564	-69.4		
611 K1FE(C1N)6-3(AO)			251. 0564		251. 0564	95.2		
612 K3FE(C1N)6(C)			329. 2604		329. 2604	-30.97		
613 K3FE(C1N)6(A)			329. 2604		329. 2604	-28.8		
FROM 3K+, FE(CN)6-3								
614 K3FE(C1N)6(500H2O)			329. 2604		-46.41			
615 K3FE(C1N)6(600H2O)			329. 2604		329. 2604	-46.34		
616 K3FE(C1N)6(800H2O)			329. 2604		329. 2604	-46.30		
617 K3FE(C1N)6(1000H2O)			329. 2604		329. 2604	-46.28		
618 K3FE(C1N)6(2000H2O)			329. 2604		329. 2604	-46.26		
619 K3FE(C1N)6(5000H2O)			329. 2604		-46.30			
620 K3FE(C1N)6(10000H2O)			329. 2604		329. 2604	-46.35		
621 K3FE(C1N)6(20000H2O)			329. 2604		329. 2604	-46.42		
622 K3FE(C1N)6(50000H2O)			329. 2604		329. 2604	-46.50		
623 K3FE(C1N)6(100000H2O)			329. 2604		329. 2604	-46.55		
624 K4FE(C1N)6(C)			368. 3624		-142.0			
625 K4FE(C1N)6(A)			368. 3624		-132.4	-104.71		
FROM 4K+, FE(CN)6-4								
626 K4FE(C1N)6(400H2O)			368. 3624		-131.71			
627 K4FE(C1N)6(500H2O)			368. 3624		-131.60			
628 K4FE(C1N)6(1000H2O)			368. 3624		-131.45			
629 K4FE(C1N)6(2000H2O)			368. 3624		-131.42			
630 K4FE(C1N)6(5000H2O)			368. 3624		-131.50			
631 K4FE(C1N)6(10000H2O)			368. 3624		-131.59			
632 K4FE(C1N)6(20000H2O)			368. 3624		-131.82			
633 K4FE(C1N)6(50000H2O)			368. 3624		-131.91			
634 K4FE(C1N)6(3H2O(C)			422. 4086		-350.5			
635 K3FE1C1(C1N)5(C)			331. 2531		331. 2531	-138.9		
636 K3FE1C1(C1N)5(A)			331. 2531		331. 2531	-134.9		
637 K3FE1C10(C1N)5(AU)			331. 2531		331. 2531	-134.5		
638 K3FE1C10(C1N)5:7/2H2O(C)			394. 3070		394. 3070	-384.0		

NBS	ALKALI	METAL	CMPD.	THERMO.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL·K				
						FORMULA	WT	DH0	DG298	H298 - H0	S298
639	K1H2FE(C1N)6(AU)					253.0724		75.0			CP298
640	K1H3FE(C1N)6(A)					254.0804		48.6			
641	K1H3FE(C1N)6(AU)					254.0804		50.0			
642	K2H1FE(C1N)6(AU)					291.1664		14.0			
643	K2H2FE(C1N)6(A)					292.1744		-11.7	21.97		101.
E44	K2H2FE(C1N)6(AU)					292.1744		-10.3			
E45	K3H1FE(C1N)6(A)					330.2684		-72.1	-42.70		115.
E46	K3H1FE(C1N)6(AU)					330.2684		-70.9			
E47	K1H2FE1C10(C1N)5(A)					255.0651		-14.0			
E48	K1H2FE1C10(C1N)5(AU)					255.0651		-13.6			
E49	K2H1FE1C10(C1N)5(A)					293.1591		-74.6			
E50	K2H1FE1C10(C1N)5(AU)					293.1591		-74.2			
E51	K1PDI1CL3(A)					251.8610		-133.7			
E52	K2PDI1CL4(C)					326.4160		-259.7			
E53	K2PDI1CL4(A)					326.4160		-245.4	-235.0		111.
E54	K2PDI1CL4(AU)					326.4160		-246.5			
E55	K2PDI1CL6(C)					397.3220		-278.0	-242.0		79.
E56	K2PDI1CL6(A)					397.3220		-264.	-238.2		114.
E57	K2PDI1CL6(AU)					397.3220		-264.			
E58	K1PDI1BR3(A)					385.2290		-116.5			
E59	K2PDI1BR4(C)					504.2400		-219.1			
E60	K2PDI1BR4(A)					504.2400		-209.4	-211.4		119.
E61	K2PDI1BR4(AU)					504.2400		-207.1			
E62	K2PDI1BRE(A)					664.0580		-215.5			
E63	K2PDI1I4(A)					692.2216		-173.4			
E64	K2PDI1I6(A)					946.0304		-176.1			
E65	K2PD(N1O2)4(A)					368.6260		-156.5			
E66	K2PD(C1N)4(A)					288.6756		15.			
E67	K2PD(C1N1S)4(A)					416.9316		-37.3			
E68	K3RH1CL6(C)					432.9290		-377.			
E69	K3RH1CL6(A)					432.9290			-383.8		
E70	K1RU1O4(A)					204.1696			-126.4		
E71	K2RU1O4(A)					243.2716			-208.0		
E72	K1PT1CL3(A)					340.5510			-122.3		
E73	K2PT1CL4(C)					415.1060			-252.6		
E74	K2PT1CL4(A)					415.1060			-240.9		89.
E75	K2PT1CL4(AU)					415.1060			-240.9		
E76	K2PT1CL6(C)					486.0120			-295.1		49.09
E77	K2PT1CL6(A)					486.0120			-282.		79.8
E78	K2PT1CL6(AU)					486.0120			-281.6		101.6

NBS ALKALI METAL CMPD.	THERMO.	PROPS.	(TN 270 SERIES) IN KCAL/MOL AND CAL/MOL·K				
			FORMULA WT	DHO	DH298	H298 - H0	S298
679 K2PT1BR4(C)			592.9300	-219.8			CP 298
680 K2PT1BR4(AU)			592.9300	-210.			
681 K2PT1BR6(C)			752.7480	-246.4			
682 K2PT1BR6(AU)			752.7480	-235.			
683 K2PT1I6(AU)			1034.7204	-172.			
684 K2(FT(N1O2)4)(C)			457.3160	64.9			
685 K2(PT1N1O2CL3)(C)			425.6585	50.4			
686 K2(PT(N1O2)2CL2)(C)			436.2110	53.3			
	CIS						
687 K2(FT(N1O2)3CL)(C)			446.7635	62.8			
688 K(PT1N1H3CL3)(C)			357.5817	-181.6			
689 K2PT(C1N)4(A)			377.3656	33.5			
690 K2IR1CL6(C)			483.1220	-281.			
691 K2IR1CL6(AU)			483.1220	-268.7			
692 K3IR1CL6(C)			522.2240	-369.			
693 K3IR1CL6(AU)			522.2240	-360.5			
694 K2OS1CL6(C)			481.1220	-285.			
695 K1MN1O4(C)			158.0376	-200.1	-176.3	41.04	
696 K1MN1O4(A)			158.0376	-189.7	-174.6	28.10	
697 K1MN1O4(14.OH20)			158.0376	-190.24			
698 K1MN1O4(200OH20)			158.0376	-190.09			
699 K1MN1O4(300H20)			158.0376	-189.99			
700 K1MN1C4(500H20)			158.0376	-189.86			
701 K1MN1O4(1000H20)			158.0376	-189.87			
702 K1MN1O4(1500H20)			158.0376	-189.75			
703 K1MN1O4(2000H20)			158.0376	-189.73			
704 K1MN1O4(5000H20)			158.0376	-189.70			
705 K1MN1O4(10000H20)			158.0376	-189.71			
706 K1MN1O4(00H20)			158.0376	-189.7			
707 K2MN1O4(A)			197.1396	-277.	-255.1	63.	
708 K1MN1CL3(A)			200.3990	-215.9			
709 K4MN(C1N)6(A)			367.4534	-108.			
710 K1RE1O4(A)			289.2996	-248.5	-233.7	72.6	
711 K2RE1CL6(A)			477.1220	-303.	-276.	109.	
712 K2CH1O4(C)			194.1976	-335.4	-309.6	47.8	
713 K2CH1O4(A)			194.1976	-331.24	-309.36	61.0	
714 K2CR1O4(17H20)			194.1976	-332.61			
715 K2CW1O4(20H20)			194.1976	-332.53			
716 K2CR1O4(25H20)			194.1976	-332.38			
717 K2CR1O4(35H20)			194.1976	-332.25			
718 K2CW1O4(45H20)			194.1976	-332.03			

NBS	ALKALI	METAL	CMPD.	THERMO.	PHOPS.	(TN	270	SERIES)	IN	KCAL/MOL	AND	CAL/MOL•K	DG298	H298	-	H0	S298	CP298
						FORMULA	WT	DHO					DH298					
719	K2CR104	(55H2O)				194.1976				-331.86								
720	K2CR104	(60H2O)				194.1976				-331.72								
721	K2CR104	(75H2O)				194.1976				-331.57								
722	K2CH104	(100H2O)				194.1976				-331.40								
723	K2CR104	(200H2O)				194.1976				-331.17								
724	K2CH104	(300H2O)				194.1976				-331.09								
725	K2CR104	(400H2O)				194.1976				-331.06								
726	K2CR104	(500H2O)				194.1976				-331.04								
727	K2CR104	(600H2O)				194.1976				-331.03								
728	K2CH104	(800H2O)				194.1976				-331.02								
729	K2CR104	(1000H2O)				194.1976				-331.03								
730	K2CH104	(2000H2O)				194.1976				-331.06								
731	K2CR104	(4000H2O)				194.1976				-331.09								
732	K2CR104	(5000H2O)				194.1976				-331.10								
733	K2CR104	(10000H2O)				194.1976				-331.12								
734	K2CR104	(20000H2O)				194.1976				-331.15								
735	K2CR104	(50000H2O)				194.1976				-331.18								
736	K2CR104	(100000H2O)				194.1976				-331.19								
737	K2CR104	(0.7H2S104+70OH2O)				194.1976				-332.31								
738	K2CR104	(1.40OH2S104+700OH2O)				194.1976				-332.59								
739	K2CR104	(3.50OH2S104+700H2O)				194.1976				-332.87								
740	K2CR104	(7.00OH2S104+700H2O)				194.1976				-333.03								
741	K2CR104	(14.00OH2S104+700H2O)				194.1976				-333.31								
742	K2CR104	(17.50OH2S104+700H2O)				194.1976				-333.47								
743	K2CH104	(35.00OH2S104+700H2O)				194.1976				-334.35								
744	K2CR104	(70.00OH2S104+700H2O)				194.1976				-336.91								
745	K2CH104	(87.50OH2S104+700H2O)				194.1976				-338.51								
746	K2CR104	(140.00OH2S104+700H2O)				194.1976				-343.69								
747	K2CR104	(175.00OH2S104+700H2O)				194.1976				-347.96								
748	K2CR207	(C)				294.1918				-492.8				-449.9				
749	K2CR207	(A)				294.1918				-476.8				-446.4				
750	K2CR207	(200H2O)				294.1918				-477.5								
751	K2CR207	(400H2O)				294.1918				-477.3								
752	K2CR207	(500H2O)				294.1918				-477.1								
753	K2CR207	(600H2O)				294.1918				-477.0								
754	K2CR207	(700H2O)				294.1918				-476.9								
755	K2CR207	(800H2O)				294.1918				-476.8								
756	K2CR207	(900H2O)				294.1918				-476.7								
757	K2CR207	(1000H2O)				294.1918				-476.6								
758	K2CR207	(1500H2O)				294.1918				-476.4								

NBS ALKALI METAL CMPD. THERMO. PROPS. (IN 270 SERIES) IN KCAL/MOL AND CAL/MOL-K

CP 298

		FORMULA WT	DH _O	DH ₂₉₈	DG ₂₉₈	H ₂₉₈ - H _O	S ₂₉₈
759	K2CH207(2000H2O)	294.1918	-4.76•2				
760	K2CR207(3000H2O)	294.1918	-4.75•8				
761	K2CF207(4000H2O)	294.1918	-4.75•6				
762	K2CH207(5000H2O)	294.1918	-4.75•5				
763	K2CR207(1000H2O)	294.1918	-4.74•2				
764	K2CH207(0.1H1CL104+1000H2O)	294.1918	-4.74•0				
765	K2CR207(0.14H1CL104+1500H2O)	294.1918	-4.73•6				
766	K2CR207(0.23H1CL104+25000H2O)	294.1918	-4.73•0				
767	K2CR207(2.80OH2S104+28000H2O)	294.1918	-4.75•86				
768	K2CH207(5.60OH2S104+28000H2O)	294.1918	-4.75•90				
769	K2CR207(14.00OH2S104+28000H2O)	294.1918	-4.75•82				
770	K2CH207(28.00OH2S104+28000H2O)	294.1918	-4.75•74				
771	K2CR207(56.00OH2S104+28000H2O)	294.1918	-4.75•60				
772	K2CR207(70.00OH2S104+28000H2O)	294.1918	-4.75•64				
773	K2CH207(140.00OH2S104+28000H2O)	294.1918	-4.75•78				
774	K2CR207(280.00H2S104+28000H2O)	294.1918	-4.78•64				
775	K2CH207(350.00H2S104+28000H2O)	294.1918	-4.80•56				
776	K2CR207(560.00H2S104+28000H2O)	294.1918	-4.86•84				
777	K2CR207(700.00H2S104+28000H2O)	294.1918	-4.93•60				
778	K2CF207(800.00H2S104+28000H2O)	294.1918	-501.00				
779	K2CR204:CR103(C)	346.1878	-6.08•				
780	K1H1CR104(A)	156.1036	-270.2				68.5
781	K3CH104F(C)	252.2980	-470.6				
782	K1CL:CR103(C)	174.5492	-248.7				
783	K1CR(S104)2(C)	283.2212	-512.				
784	K1CR(S1C4)2(AU)	283.2212	-562•0				
785	K1CR(S104)2:H2O(C)	301.2366	-592•				
786	K1CH(S104)2:H2O(C)	319.2520	-669•				
787	K1CR(S104)2:6H2O(C)	391.3136	-966•0				
788	K1CR(S104)2:12H2O(C)	499.4060	-1389•2				
789	K1N1H4CF104(C)	173.1343	-307•8				
790	K1N1H4CR104(AU)	173.1343	-302•5				
791	K2MC104(C)	238.1416	-358•2				
792	K2MC104(A)	238.1416	-359•1				55.5
793	K2MC104(AU)	238.1416	-358•9				
794	K2W104(A)	326.0516	-377•7				
795	KEH1W6O21(A)	1635.6054	-1697•2				
796	K1W1CL6(C)	435.6700	-235•4				
797	K2W1CL7(C)	510.2250	-338•4				
798	K1V1O3(C)	138.0422	-276•0				

NBS	ALKALI	METAL	CMPD.	THERMO.	PROPS.	(TN 270 SERIES)	IN KCAL/MOL AND CAL/MOL.K			
						FORMULA	WT	DHO	DH298	DG298
799	K1V1C3(A)					138.0422		-272.6	-255.0	37.
800	K3V1O4(A)					232.2456			-418.0	
801	K1H2V1O4(A)					156.0576		-340.9	-311.7	54.
802	K1H3V2O7(A)					256.0058			-513.2	
803	K3H1V2O7(A)					332.1938			-631.5	
804	K4H2V1O028(A)					1115.8272		-2380.	-2117.	176.
805	K5H1V1O028(A)					1153.9212			-2180.	
806	K1TA1F6(A)					334.0404			-276.9	
807	K2TA1F7(A)					392.1408			-415.6	
808	K1MG1FE(C1N)6(A)					275.3684			-5.9	
809	K2MG1FE(C1N)6(A)					314.4704			-83.2	
810	K1CA1FE(C1N)6(A)					291.1364			-29.6	
811	K2CA1FE(C1N)6(A)					330.2384			-106.8	
812	K1SH1FE(C1N)6(A)					338.6764			-31.0	

CP298

		IN KCAL/MOL AND CAL/MOL.K	CP298
1	PR(CS)	3 / 76 DH298	DG298
2	FB(G)	85.4678 0 0 0	H298 - HO 1.790 18.35 ♦-0.005 ♦-0.07
3	PR+(C)	85.4678 1C.639 1C.37 12.69 ♦-0.050 ♦-0.05 ♦-0.05	40.626 4.968 ♦-0.005 ♦-0.005
4	PR+2(G)	85.4678 745.10 747.76 ♦-0.06 ♦-0.06	
5	PR+3(G)	85.4678 1660.0 1664.1 ♦-2.0 ♦-2.0	
6	PR+(A)	85.4678 -60.03 -67.87 ♦-0.03 ♦-0.03	29.04 ♦-0.05
7	PR1E2(C)	117.4666 -66.6 ♦-0.5	
8	PR2E(C)	186.9350 -81. ♦-3.	
9	PR2E(G)	186.9350 -12. ♦-5.	
10	PR2E2(C)	202.9344 -112.8 ♦-2.5	
11	PR1H(C)	86.4758 -12.5 ♦-0.3	
12	PR1E1H(C)	102.4752 -99.95 ♦-0.15	
13	PR1E1H(G)	102.4752 -56. ♦-2.	-57. ♦-2.
14	PR1E1H(A)	102.4752 -115.00 -105.46 ♦-0.03 ♦-0.03	26.47 ♦-0.05
15	PR1E1H:B29(C)	120.4906 -178.98 ♦-0.15	-157. ♦-12.
16	PR1E1H:2H29(C)	138.5060 -251.73 ♦-0.15	
17	(PR1E1H)2(G)	204.9504	

NPS PURIDIUM CMPDS THERM⁹ PREPS (TN270 SERIES) IN KCAL/MOL AND CAL/MOL.K
 FORMULA WT DH₀ DG298 H298 - H₀ S298 CP298

18 RB1F(C) 104.4662 -133.3 -133.3

19 RB1F(G)
 SPECTRAL DATA FROM RESEN.
 FROM AKISHIN AND RAMBINI.
 KHIM, 4, 768 (1958) 104.4662 -78.65 -79.2 +0.2
 ZHUR. NEORG. +1.12 +1.1 +1.5 +0.15 +0.15

20 RB1F(A) 104.4662 -139.53 -134.51 +0.08 +0.08 +0.3

21 RB1F(100H2 Θ) 104.4662 -139.35 -139.35 +0.08

22 RB1F(150H2 Θ) 104.4662 -139.40 -139.40 +0.08

23 RB1F(200H2 Θ) 104.4662 -139.408 -139.408 +0.080

24 RB1F(300H2 Θ) 104.4662 -139.419 -139.419 +0.080

25 RB1F(400H2 Θ) 104.4662 -139.428 -139.428 +0.080

26 RB1F(500H2 Θ) 104.4662 -139.435 -139.435 +0.080

27 RB1F(600H2 Θ) 104.4662 -139.441 -139.441 +0.080

28 RB1F(800H2 Θ) 104.4662 -139.450 -139.450 +0.080

29 RB1F(1000H2 Θ) 104.4662 -139.456 -139.456 +0.080

30 RB1F(1500H2 Θ) 104.4662 -139.467 -139.467 +0.080

31 RB1F(2000H2 Θ) 104.4662 -139.473 -139.473 +0.080

32 RB1F(3000H2 Θ) 104.4662 -139.482 -139.482 +0.080

33 RB1F(5000H2 Θ) 104.4662 -139.491 -139.491 +0.080

34 RB1F(10000H2 Θ) 104.4662 -139.501 -139.501 +0.080

35 RB1F(20000H2 Θ) 104.4662 -139.505 -139.505 +0.080

36 RB1F(50000H2 Θ) 104.4662 -139.517 -139.517 +0.080

37 RB1F(100000H2 Θ) 104.4662 -139.520 -139.520 +0.080

38 RB1F(1.5H2 Θ (C)) 131.4893 +0.3 +0.3

NBS RUBIDIUM CMPDS THERMO PROPS (IN 270 SERIES) IN KCAL/MOL AND CAL/MOL.K				3/76			
FORMULA	WT	D ₁₀₀	DH ₂₉₈	DG ₂₉₈	H ₂₉₈	H ₀	S ₂₉₈
3C RB1F(H1C181N1H2:S) IN FERMAMIDE	104.4662	-138.57 ♦-0.20					CP298
40 RB2F2(G)	208.9324	-204. ♦-5.					
41 RB1H2(C) THERMAL FUNCTIONS FROM BURNEY, G. A.; AND WESTRUM, E. F., JR.; J. PHYS. CHEM. 65, 349 (1961)	124.4726	-219.52 ♦-0.21	-220.5 ♦-0.2	-204.5 ♦-0.2	3.932 ♦-0.004	28.7 ♦-0.0	18.97 ♦-0.02
42 RB1CL(C) THERMAL FUNCTIONS FROM PAUKEV, I. E.: AND KHPIPILOVICH, L. V.; ZHUR. FIZ. KHM. 43, 2678 (1969), REINTERPATED AT NBS	120.9208	-104.080 ♦-0.051	-104.05 ♦-0.05	-97.47 ♦-0.05	2.917 ♦-0.004	22.92 ♦-0.04	12.52 ♦-0.04
43 RB1CL(G) SPECTRAL DATA FROM ROSEN, BOND LENGTH FROM AKISHIN AND RAMBIDI, ZHUR. NEORG. KHM., 4, 768 (1959)	120.9208	-55.11 ♦-1.12	-55.6 ♦-1.1	-60.0 ♦-1.2	2.40 ♦-0.15	59.6 ♦-1.0	8.78 ♦-0.15
44 RB1CL(A)	120.9208	-99.98 ♦-0.03	-99.24 ♦-0.03	-99.24 ♦-0.03		42.54 ♦-0.05	
45 RB1CL(50H1C161H) IN FORMIC ACID	120.9208	-103.1 ♦-0.1					
46 RB1CL(H1C161H2:S) IN FERMAMIDE	120.9208	-103.35 ♦-0.05					
47 RB1CL(H1C161N1H1C1H3:S) IN N-METHYLFOPMAMIDE	120.9208	-103.21 ♦-0.05					
48 RB1CL(CAH8H2:S) IN 20 PERCENT DIOXANE-WATER	120.9208	-100.38 ♦-0.05					
49 RB1CL(C1H361H:U) IN METHANOL	120.9208	-101.86 ♦-0.08					
50 (RB1CL)2(G)	241.8416	-150. ♦-5.					
51 RB1CL163(C) S AND CP FROM KELLEY, K. K.; AND KING, E. G.; US PUR. MINES BULL. 592, 149P (1961)	168.9190	-95.1 ♦-0.3	-70.6 ♦-0.3			36.3 ♦-0.4	24.66 ♦-0.05
52 RB1CL163(A)	168.9184	-83.7 ♦-0.2	-68.7 ♦-0.2				67.8 ♦-0.5
53 RB1CL164(C)	184.9184	-104.50 ♦-0.10	-73.54 ♦-0.10				39.2 ♦-0.5
54 RB1CL164(A)	184.9184	-90.94 ♦-0.08	-69.93 ♦-0.08				72.5 ♦-0.5
55 RB1CL164(C4H8S1H2)	184.9184	-102.25 ♦-0.25					
56 RB1CL164(H1C161N1H2:S) IN FERMAMIDE	184.9184	-100.36 ♦-0.13					

NBS PUBLISHED COMPOUNDS THERMOPROPS (TN270 SERIES) IN KCAL/MOL AND CAL/MOL.K									
FORMULA WT	DHO	DH298	DG298	H298	H0	S298			CP298
57 RB1CL1θ4(H1C1θ1N1H1C1H3 : S) IN N-METHYLFORMAMIDE	184.9184	-101.70 + -0.13							
58 RB1CL1θ4(H1C1θ1N(C1H3)2 : S) IN N,N-DIMETHYLFORMAMIDE	184.9184	-106.35 + -0.13							
59 RB1BR(C) THERMAL FUNCTIONS FROM CLUSIUS, K.; GELDMANN, J.; AND PERLICK, A.; Z. NATURFORSCH. 4A, 424 (1949), REINTEGRATED AT NBS	165.3768	-92.714 + -0.052	-94.31 + -0.05	-91.24 + -0.05	3.124 + -0.005	26.28 + -0.05	12.63 + -0.01		
60 RB1BR(G) SPECTRAL DATA FROM ROSEN. BOND LENGTH FROM AKISHIN AND RAMBIDI, ZHUR. NEORG. KHIM., 4, 768 (1959)	165.3768	-42.15 + -1.02	-44.4 + -1.0	-52.1 + -1.3	2.47 + -0.15	62.4 + -1.0	8.87 + -0.15		
61 RB1BR(A)	165.3768	-89.08 + -0.08	-92.72 + -0.08			48.74 + -0.05			
62 RB1BR(H1C1θ1N1H2 : S) IN FORMAMIDE	165.3768	-93.56 + -0.25							
63 RB1BR3(C)	325.1948	-100.0 + -1.3							
64 RB1BR1θ3(C)	213.3750	-91.72 + -0.50	-70.52 + -0.50			38.9 + -0.9			
65 RB1BR1θ3(A)	213.3750	-80.0 + -0.4	-67.5 + -0.4			68.0 + -0.8			
66 RB1BR1CL2(C)	236.2828	-116.0 + -1.5	-99.7 + -0.5			35.1 + -2.0			
67 RB1BR2CL(C)	280.7388	-112.5 + -1.0	-99.1 + -0.5			36.6 + -2.0			
68 RB1I(C) THERMAL FUNCTIONS FROM CLUSIUS, K.; GELDMANN, J.; AND PERLICK, A.; Z. NATURFORSCH. 4A, 424 (1949), REINTEGRATED AT NBS	212.3722	-79.593 + -0.123	-79.77 + -0.12	-78.60 + -0.07	3.190 + -0.010	28.30 + -0.05	12.71 + -0.05		
69 RB1I(G) SPECTRAL DATA FROM ROSEN. BOND LENGTH FROM AKISHIN AND RAMBIDI, ZHUR. NEORG. KHIM., 4, 768 (1959)	212.3722	-31.74 + -1.12	-32.6 + -1.1	-42.2 + -1.3	2.51 + -0.15	64.3 + -1.0	8.92 + -0.15		
70 RB1I(A)	212.3722								
71 PB1I((1000N1H2C1H2C1H2N1H2) ETHYLENEDIAMINE	212.3722								
72 RB1I(H1C1θ1N1H2 : S) IN FORMAMIDE	212.3722								
73 RB1I(H1C1θ1N1H1C1H3 : S) IN N-METHYLFORMAMIDE	212.3722								
74 RB1I(H1C1θ1N1(C1H3)2 : S) IN N,N-DIMETHYLFORMAMIDE	212.3722								

			KCAL/MOL	CAL/MOL.K							
			WT	DHO							
75	RBI1(C1H3C1=O1N1H1C1H3:S2)		212.3722		-80.08						CP298
	IN N-METHYLACETAMIDE				+ -0.15						
76	RBI1(C1H3C1=O1N1H1C1H3:S2)		212.3722		-81.60						
	COMPOUND OUT OF ORDER				+ -0.15						
77	RBI1(C)Nitrile		466.1810		-82.7	-81.0					
	IN ACETONITRILE				+ -0.3	+ -0.1					
78	RBI1103(C)		260.3704		-101.9						
					+ -0.7						
79	RBI1103(A)		260.3704		-112.9	-98.5					
					+ -0.5	+ -0.5					
80	RBI111CL2(C)		283.2782		-106.1						
					+ -1.0						
81	RBI111CL4(C)		354.1842		-129.4	-105.3					
					+ -0.5	+ -0.5					
82	RBI111BR2(C)		372.1902		-96.0						
					+ -0.5						
83	RBI111BR1CL(C)		327.7342		-102.3						
					+ -0.5						
84	RB2S(C)		202.9996		-86.2						
					+ -1.0						
85	RB2S(500H2θ)		202.9996		-111.0						
					+ -1.0						
86	RB1S2θ-(Aθ)		277.5910		-378.7	-334.9					
					+ -0.7	+ -0.7					
87	RB2S1θ4(C)		266.9972	-340.795	-343.12	-314.76					
	THERMAL FUNCTIONS FROM PAUKEV, I. E.;			+ -0.075	+ -0.07	+ -0.07					
	AND LAURENT'eva.										
	4.2°, 184.2 (1968)										
88	RB2S1θ4(G)		266.9972		-259.						
89	RB2S1θ4(A)		266.9972		+ -6.						
					+ -0.05	+ -0.05					
90	RB2S1θ4(500H2θ)		266.9972		-337.161						
					+ -0.050						
91	RB2S1θ4(800H2θ)		266.9972		-337.156						
					+ -0.050						
92	RB2S1θ4(1000H2θ)		266.9972		-337.157						
					+ -0.050						
93	RB2S1θ4(1500H2θ)		266.9972		-337.164						
					+ -0.050						
94	RB2S1θ4(2000H2θ)		266.9972		-337.170						
					+ -0.050						
95	RB2S1θ4(3000H2θ)		266.9972		-337.183						
					+ -0.050						

NBS RUBIDIUM CMPDS THERMD PREPS (TN270 SERIES) IN KCAL/MOL AND CAL/MOL.K	3/76	DH298	H298 - HO	S298	CP298
FORMULA	WT	DHO			
96 RB2S104(5000H2O)	266.9972	-337.204 + -0.050			
97 RB2S104(1000OH2O)	266.9972	-337.242 + -0.050			
98 RB2S104(2000OH2O)	266.9972	-337.276 + -0.050			
99 RB2S104(5000OH2O)	266.9972	-337.310 + -0.050			
100 RB2S104(10000OH2O)	266.9972	-337.329 + -0.050			
101 RB2S104(20000OH2O)	266.9972	-337.343 + -0.050			
102 RB2S104(5000OH2O)	266.9972	-337.356 + -0.050			
103 RB1S104(C) CP FRM TEICHERT, W.; Z. ANORG. ALLGEM. CHEM. 247, 113 (1941)	118.5398	-53.8 + -1.1			14.6 + -1.0
104 RB1S104(1000H2O)	118.5398	-64.5 + -1.0			
105 RB1H1S104(C)	182.5374	-277.0 + -0.5			
106 RB1S102F(C)	168.5290	-225.7 + -0.7			
107 RB1I:3S102(C)	404.5606	-324.0 + -1.0			
108 RB2SE103(C)	297.8936	-225.7 + -0.5			
109 RB2SE104(C)	313.8932	-266.3 + -0.1			
110 RB1H1SE(C) CP FRM TEICHERT, W.; Z. ANORG. ALLGEM. CHEM. 247, 113 (1941)	165.4358	-57.2 + -0.5			15.7 + -1.0
111 RB2TE1BR6(C)	777.9896	-401.6 + -0.5			
112 RB1N3(C)	127.4879	-0.9 + -0.1			
113 RB1N103(C) CP FRM MUSTAJEKI, A., ANN. ACAD. SCI. FINNICAЕ [A] VI, № 9, 16P (1958)	147.4727	-118.32 + -0.05	-94.61 + -0.05		24.4 + -1.0
114 RB1N103(A)	147.4727	-109.59 + -0.05	-94.48 + -0.05		64.0 + -0.1
115 RB1N1H2(C)	101.4905	-27.0 + -0.5			
116 RB1F2P104(C)	182.4552	-373.39 + -0.10			

	NRS PURIDUM CMPPDS THERM. PRAPS (TN270 SERIES) IN KCAL/MOL AND CAL./MOL.K			3/76			
	FORMULA	WT	DH ₀	DH ₂₉₈	H298 - H0	S298	CP298
117 RB1PIF6(C)	THEMAL FUNCTIONS FROM STAVELEY, L. A. K.; GREY, N. R.; AND LAYZELL, M. J.; Z. NATURFORSCH. 18A, 148 (1963)	230.4320	-562.8 + -1.0	-526.9 + -1.0	53.02 + -0.15	35.4 + -0.1	
118 RR1SP(C)		207.2178	-23.9 + -1.5	-24.5 + -1.5			
119 RB1SP2(C)		328.5678					
120 RB3SB(C)		378.1534	-41.5 + -1.0	-74.1 + -5.0			
121 RB3SB7(C)		1108.6534					
122 RB5SB4(C)		914.3350	-106.1 + -4.0				
123 SB1C13:3RB1C1(C)		590.8714	-410.2 + -0.8	-884.6 + -2.0			
124 (3RB1RR:3SB1BR3)(C)		2242.0686					
125 RB2C163(C)	TMRAL FUNCTIONS FROM PAUKOV, I. E.; KRIVLEVICH, L. M.; AND LUK'YANOVA, I. G.; ZHUR. FIZ. KHIM. 45, 2451 (1971)	230.9450	-270.41 + -0.22	-251.1 + -0.5	5.851 + -0.010	43.34 + -0.10	
126 RF2C163(A)		230.9450					
127 RB2C163(5.76826)		230.9450	-280.0 + -0.5				
128 RB2C163:H26(C)		248.5604	-346.1 + -0.2				
129 RB2C163:1.5H26(C)		257.9681	-383.4 + -0.2				
130 RB2C163:3.5H26(C)		293.5985	-523.6 + -0.5				
131 PH1C163(C)		146.4852	-230.2 + -0.4	-206.3 + -0.7			
132 RB1H1C163(A)		146.4852	-225.42 + -0.10	-208.13 + -0.10			
133 (3RF2C163:2RP1H1C163:4.5H26)(C)		1066.8747	-1606.2 + -0.8				
134 RB1C1(N)		111.4857					
135 RB2SIIIF6(C)	TMRAL FUNCTIONS FROM SUGISAKI, M.; MATSUO, T.; SUGA, H.; AND SEKI, S.; BULL. CHEM. SOC. JAPAN 41, 1747 (1968)	313.0120	-695.5 + -2.5	33.67 + -0.05	16.20 + -0.02		
136 RB2GE1C16(C)		456.2436	-350.0 + -1.5	-310.8 + -1.0	72.5 + -2.5		

NBS RUBIDIUM CMPS THERM PROPS (TN270 SERIES) IN KCAL/MOL AND CAL/MOL.K FORMULA WT DH0	DH298	DG298	H298 - H0	S298	CP298
137 PB1SN1CL6(C) S AND CP FROM MORFEE, R. G. S.; STALEVELY, L. A. K.; WALTERS, S. T.; AND WIGLEY, D. L.; PHYS. CHEM. SOLIDS. 13, 132 (1960)	502.3436	-364.0 + -1.0	-3228.6 + -1.0	90.25 + -0.05	54.27 + -0.05
138 RB2SN1BR6(C)	769.0796	106.4 + -0.1	54.54 + -0.05		
140 PB1I2:2PB1I:4H2O(C)	957.8048	-488.9 + -0.5	3.181 + -0.010	22.54 + -0.06	17.7 + -0.1
141 RB1B162(C) THERMAL FUNCTIONS FROM PAUKEV, I. E.; KHRIKAVICH, L. M.; AND POPEN, A. P.; ZUR. FIZ. KHIM. 45, 1295 (1971)	128.2776				
142 RB1B1F4(C) CP FROM DWERKIN, A. S.; AND BREDIG, M. A.; J. CHEM. ENG. DATA 15, 505 (1970)	172.2724	-449.3 + -2.5			22.7 + -0.3
143 PB1B1CL4(C)	238.0908	-224.7 + -0.5			
144 RB1R1(CL164)4(C)	494.0812	-185.7 + -1.3			
145 RB1AL1(SE164)2:12H2O(C)	614.5493	+1303.3 + -2.6			
146 RB2ZN1CL4(C)	378.1176	-319.3 + -0.2			
147 RB2ZN1BR4(C)	555.9416	-277.14 + -0.03			
148 RR1CL:2N1S164(C)	282.3524	-346.4 + -0.1			
149 CU1CL2:2RB1CL(C)	376.2876	-266.3 + -0.2			
150 CU1CL2:2PB1CL:4H2O(C)	448.3492	-547.2 + -0.2			
151 PB1AG4I5(C) H+BO AND CP FROM TOPAL, L. E.; AND GWENS, B. B.; J. PHYS. CHEM. 72, 2106 (1968)	1151.46c8	-138.82 + -1.05	-143.0 + -1.0	17.092 + -0.10	149. + -1.
152 RB2AG1I3(C)	659.5188	-176.2 + -0.4	-175.2 + -0.2	85.	
153 RB1NI1CL3(C)	250.5368	-182.7 + -0.2			
154 PR1CP1CL3(C)	250.7600	-184.4 + -0.3			

NBS RUBIDIUM CMPDS THERM. PREPS (TN270 SERIES) IN KCAL/MOL AND CAL/MOL.K			3/76				
	FOPMULA WT	DHO	DR298	DG298	H298	- H0	S298
155 RB2C ⁶ 1CL4(C)	371.6808				=288.3		CP298
156 RB7C ⁶ 1CL5(C)	492.6016				⁺ -0.3		
157 PR1FE1CL3(C)	247.6738				-394.0		
158 PR2FE1CL4(C)	368.5946				⁺ -0.3		
159 RB2PT1CL4(C)	507.8376				-190.8		
160 RB2PT1CL6(C)	578.7436				⁺ -0.2		
161 RB1[PT1N1H3CL3](C)	403.9475				-294.6		
162 RB2TR1CL6(C)	575.8536				⁺ -0.2		
163 RB1NN1CL3(C)	246.7648				-255.4		
164 RB1RF1H4(C)	335.5654				⁺ -0.3		
165 RB2CR1H4(C)	286.9292				-299.		
166 RB3CP1e4F(C)	391.3954				⁺ -5.		
167 RB3CR1CL6(C)	521.1174				-267.		
168 RB3CP2CL9(C)	679.4724				⁺ -3.		
169 RB3V1CL6(C)	520.0634				-225.		
170 RB3V2CL9(C)	677.3644				⁺ -2.		
171 RB1NB1e3(C)	226.3720				-182.4		
172 RB1NB1CL6(C)	391.0918				⁺ -0.3		
173 RB2NP1H1CL5(C)	457.1060				-160.		
174 RB1TA1CL6(C)	479.1338				⁺ -6.		
175 RB1TT1CI3(C)	239.7268				-326.9		
176 RB2TT1CL4(C)	360.6476				⁺ -0.8		
					-250.1		
					⁺ -1.5		
					-358.0		
					⁺ -2.0		

NBS RUBIDIUM CMPDS THERM PAPPS (TN270 SERIES) IN KCAL/MOL AND CAL/MOL.K
 FORMULA WT DHO DG298 H298 - H0 S298
 DH298

177 RB2T11CL6(C)	431.5536	-425. + -3.
178 RB2T11BR6(C)	698.2896	-359.7 + -0.8
179 RB3T11BR6(C)	783.7574	-425. + -0.7
180 RB3T12BR6(C)	1071.3844	-563.5 + -0.8
181 RB1GD(FF(C1N)E)(C)	454.6722	-75.9 + -1.5
182 RB1CE(FF(C1N)E):2H26(C)	473.5730	-192.8 + -1.0
183 TH1CL4:2RB1CL(C)	615.6917	-512.0 + -1.0
184 TH1CL4:2RB1CL:9H26(C)	777.8303	-1154. + -1.8
185 TH1CL4:4RB1CL(C)	857.5333	-728.0 + -1.5
186 RB2MG(SE164)2(6400H28)	481.1628	-517.38 + -1.20
187 RB2MG(SE164)2:6H26(C)	589.2552	-937.6 + -1.2
188 RB1CA1CL3(C)	231.9068	-299.5 + -0.2
189 NA1RB2CP1CL6(C)	458.6394	-456.5 + -0.5
190 KRB1CL2(C)	195.4758	-208.87 + -0.05

TOTAL COMPOUNDS 191

U.S. DEPT. OF COMM. BIBLIOGRAPHIC DATA SHEET		1. PUBLICATION OR REPORT NO. NBSIR 76-1034	2. Gov't Accession No.	3. Recipient's Accession No.
4. TITLE AND SUBTITLE CHEMICAL THERMODYNAMIC PROPERTIES OF COMPOUNDS OF SODIUM, POTASSIUM AND RUBIDIUM: AN INTERIM TABULATION OF SELECTED VALUES		5. Publication Date April 1976		
		6. Performing Organization Code		
7. AUTHOR(S) D. D. Wagman, W. H. Evans, V. B. Parker and R. H. Schumm		8. Performing Organ. Report No. NBSIR 76-1034		
9. PERFORMING ORGANIZATION NAME AND ADDRESS NATIONAL BUREAU OF STANDARDS DEPARTMENT OF COMMERCE WASHINGTON, D.C. 20234		10. Project/Task/Work Unit No. 3165151		
		11. Contract/Grant No.		
12. Sponsoring Organization Name and Complete Address (Street, City, State, ZIP) National Bureau of Standards Department of Commerce Washington, DC 20234		13. Type of Report & Period Covered Interim		
		14. Sponsoring Agency Code		
15. SUPPLEMENTARY NOTES				
16. ABSTRACT (A 200-word or less factual summary of most significant information. If document includes a significant bibliography or literature survey, mention it here.) Selected values are given for the thermochemical properties of the more common compounds of sodium and potassium. A more extensive set of selections is provided for rubidium compounds. The properties included, where data are available, are enthalpy of formation at 0 K and 298.15 K, $\Delta H_f(0)$ and $\Delta H_f(298)$, Gibbs energy of formation, entropy and heat capacity at 298.15 K, $\Delta G_f(298)$, $S(298)$ and $C_p(298)$ and the enthalpy difference between 0 K and 298.15 K, $H(298)-H(0)$. The values are consistent with the tables issued earlier in the NBS Technical Note 270 series.				
17. KEY WORDS (six to twelve entries; alphabetical order, capitalize only the first letter of the first key word unless a proper name; separated by semicolons) Enthalpy; entropy; Gibbs energy; heat capacity; potassium compounds; rubidium compounds; sodium compounds; standard reference data; thermochemical tables.				
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